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Migration of Low Molecular Weight Additives in Polyolefins and Copolymers

U.S. DEPARTMENT OF COMMERCE

National Bureau of Standards

- Center for Materials Science
Polymer Science and Standards Division
Washington, DC 20234

Final Project Report

Issued March 1982

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Prepared for
Bureau of Foods
Food and Drug Administration
Washington, DC 20201

Errata

Page III-6 Add the following after equation (1):

where $\alpha = M_{S\infty}/M_{P\infty} = V_S/KV_P$, $K = C_{P\infty}/C_{S\infty}$ and $T = Dt/\lambda^2$.

Tables IV-1.1 through III-7.3 Should be placed after page III-2.

Table III-4.1 "Triglycerides" in the title should read "Triglycerides".

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U.S. DEPARTMENT OF COMMERCE, Malcolm Baldrige, Secretary
NATIONAL BUREAU OF STANDARDS, Ernest Ambler, Director



Abstract

Food packaging is an essential part of modern life. Any substances that migrate from packaging material into foods are viewed as indirect food additives. In addition to toxicological knowledge, it is important to know the amount of such indirect food additives expected to be present in the food during storage and processing. This program, sponsored by the Bureau of Foods of the Food and Drug Administration, is designed to provide theoretical models, a reliable data base and methodology for studying migration phenomena and can provide reasonable worst-case estimates for the concentrations of the indirect additives in foods. In this final report, we present the results of approximately 250 completed migration experiments based on radiotracer techniques on the migration of low molecular weight hydrocarbons and antioxidants from polyethylene, polypropylene, and ethylene-vinyl acetate copolymers. Results of a study for the determination of relative diffusion coefficients of several probe molecules in the polyethylene melt by inverse gas chromatography are also presented. Based on these studies, ethanol appears to be a far more reasonable food-oil simulating solvent than n-heptane. Other pure or mixed triglycerides may also be considered as food-oil simulating solvents, however they may pose the same analytical difficulties associated with the use of the food oils themselves.

Keywords: antioxidants, diffusion, ethylene-vinyl acetate copolymers, food additives, food packaging, inverse gas chromatography, migration, oligomers, polyethylene, polypropylene, radiotracer.

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I. Introduction

As in the proceeding decades, most foods available today are sold in packaged forms rather than in bulk for reasons of preservation, convenience and ease of distribution. The Bureau of Foods of the Food and Drug Administration has regulatory responsibility for the use of packaging materials in contact with food. Any substances that migrate from the food packaging materials into foods are considered to be indirect food additives. The regulatory decisions on the types of packaging materials suitable for use with various types of foods are based on: the indirect food additives that can be expected to migrate from the package into the food, ones knowledge of toxicology and the expected amount of migration during storage and usage. Since it is impossible to test every conceivable combination of polymer-migrant-food, this program at the National Bureau of Standards, sponsored by the Food and Drug Administration, is aimed at providing sound technical suggestions and reliable data to the Food and Drug Administration, and recommending theoretical or empirical estimation and extrapolation methods to yield at least a reasonable worst-case estimates for regulatory purposes. However, this report represents only the technical opinions of the authors as representatives of the National Bureau of Standards and should not be taken in any way as containing regulatory decisions or recommendations of the Bureau of Foods.

A large number of kinetic experiments were performed involving the migration of radioactive ¹⁴C-labeled paraffinic oligomers (n-octadecane and n-dotriacontane) and an antioxidant (butylated-hydroxytoluene) from two different polyethylenes (high density or linear polyethylene and low density or branched polyethylene), isotactic polypropylene and two (ethylene-vinyl acetate) copolymers of relatively low vinyl acetate

content into various solvents including a variety of triglycerides, alcohols, hydrocarbons, and water at temperatures from room temperature to 60°C.

Among the correlations for various parameters, it was found that either the pure triglycerides (e.g. tributyrin and trioctanoin), mixed triglycerides, or anhydrous ethanol may successfully be employed as food-oil simulant in all cases studied. The degree of accelerating action of extracting migrants by n-heptane over that by the food oils is quantified.

Future works in the additive migration program should be mainly in the area of migration of additives from glassy polymers, where the diffusion coefficients are many orders of magnitude slower than those for the amorphous or semi-crystalline polymers above their glass transition temperatures.

Inverse gas chromatography (IGC) studies have been used to determine the relative diffusion coefficients of oligomers and antioxidants in polymers at temperatures above the glass transition region and in the supercooled-liquid or liquid phase. The determinations of the relative diffusion coefficients of several probe molecules in polyethylene are illustrated.

II. Experimental Details of Extraction Experiments

All scheduled experiments on the migration of additives from polyolefins (polyethylenes and polypropylenes) are now completed. In this report we present a complete summary of the polyolefin work, most of which has been reported in previous NBSIRs [1-6] on this subject, therefore, other than necessary descriptions to make this report self-explanatory, detailed discussions on each sub-category will not be repeated here.

As the copolymer work was started in mid-year and was just completed recently, rather detailed descriptions and discussions of it will be given in this report.

Migrations of low molecular weight additives moving from polymeric materials into surrounding liquid media under well stirred conditions were measured in this laboratory for the following combinations of base polymer, migrant, migrant concentration, solvent and temperature:

Polymers	Linear Polyethylene (LPE), SRM 1475 Branched Polyethylene (BPE), SRM 1476 Polypropylene (PP) Ethylene-5% Vinyl Acetate Copolymer Ethylene-13% Vinyl Acetate Copolymer
Migrants	n-Octadecane, n-C ₁₈ H ₃₈ n-Dotriacontane, n-C ₃₂ H ₆₆ BHT or 3,5-di-t-butyl-4-hydroxytoluene
Migrant Concentration	From 30 ppm to 10%
Solvents	Accelerating solvents: n-Heptane, n-Octadecane Fat and Oil Simulating Solvents: Corn Oil, Ethanol, n-Octanol, Tributyrin, Trioctanoin, HB307 Aqueous Solvents: Water, Ethanol-Water Mixtures
Temperatures	24, 30, 40, 60°C

The migrants are radiolabeled by carbon-14. The amount of radioactivity is determined by liquid scintillation counting techniques with detection limit less than 10 pCi.

Materials

The characteristics of the base polymers are listed in Table II-1 and II-2 and those of the base radioactive-labeled migrants are listed in Table II-3.

Sample Plaque Preparation

The following procedure for the mixing of additives in the polymer stock and the molding of the sample plaques was chosen. A large quantity of polyethylene powder stock was prepared from either the National Bureau of Standards-Standard Reference Material (NBS-SRM) 1475 or 1476 pellets by first dissolving it in hot toluene or xylene. Most of the polyethylene precipitates out upon cooling. The precipitate, together with the residue obtained by evaporating the solvent, was dried in a vacuum oven to remove the last trace of solvent.

A specific amount of labeled additive dissolved in a highly volatile solvent is mixed with a quantity of the polyethylene powder stock. The mixture, together with a number of glass beads which act as a ball mill, is then evaporated to dryness in a rotary evaporator under reduced pressure at relatively low temperatures. Further drying is carried out in a vacuum oven.

The mixture is then compression molded in a hydraulic press operated at about 180°C for oligomers and 165°C for BHT. Plaques of 125 mm x 125 mm or less are molded with brass or stainless steel shim stocks of appropriate thickness sandwiched between two sheets of Teflon or Teflon coated plates. Teflon surfaces are used to allow easy removal of the sample plaques without the use of, and subsequent contamination by, mold-release agents.

The isotactic polypropylene and (ethylene-vinyl acetate) copolymers were received in granular form from the suppliers. Therefore, they were used directly to form slurries with the migrants in solution as described above.

Special problems are encountered in the preparation of sample plaques with BHT as an additive. A much longer time is required to mix BHT with polymeric powder in a rotating flask with glass beads acting as a ball mill. Apparently BHT is relatively insoluble in the polymer even in the molten state, therefore, sample plaques made from such insufficiently mixed batches of powder mixtures yield autoradiographs of distinctively sharp regions of streaks and patches. Each molding at 185°C seems to "fix" an additional 15-20% of the available BHT, i.e. only 80% of the available BHT before the last molding may be extracted exhaustively even by n-heptane. Only 20% of original amount of BHT can be extracted from plaques remolded five times at 185°C. The residual BHT is combined with the polymer. This residual radioactivity stays in the polymeric precipitate, even after the plaques were dissolved in toluene at high temperatures. However, molding of well mixed batches of BHT and polymeric powder at 165°C seems to result in rather uniformly distributed sample plaques with less than 5% of BHT being combined with the polymer. The final fraction extracted from the original loading, M_f/M_0 , by n-heptane, as listed in Tables in Section III, gives an indication of the unreacted fraction of BHT in those samples. Therefore the original loading M_0 , is irrelevant and should be replaced by the remaining unreacted amount, M_f , for kinetic and partition calculations. There are further complications due to the volatility of BHT or its degraded components as detailed later in Section II.

Experimental Methods

Two extraction methods were used: (1) continuous extraction into a limited solvent volume and (2) discrete extraction into a simulated infinite solvent volume.

In method (1) an extraction vial of 25 ml volume with a Teflon valved cap is used. The 10 to 15 ml of solvent, in the vial will only contact glass walls and Teflon surfaces during normal experimental processes. A silicone plug is situated above the valve. A small area of the silicone rubber, less than 1 mm in diameter which is used as a septum for the hypodermic needle, is exposed to the solvent vapor. The polymer sample was sometimes surrounded by a nichrome or stainless steel screen to prevent it from sticking to another sample or to the walls, if the sample has lower density than the solvent.

The total amount extracted, at time t , M_t , is the sum of the amount of migrant in the solution at time t plus that were removed in previous aliquots,

$$M_t = C_{st} W_{st} + \sum_{i=1}^{t-1} C_{si} W_{ai}$$

where C_s , W_s and W_a represent the concentration of the migrant, total weight of the solution (including that of the aliquot for the determination of C_{st}) and the weight of the aliquot, respectively. At equilibrium the partition coefficient is estimated as the ratio of the concentrations in solvent and in polymer at equilibrium,

$$k = \frac{C_{s\infty}}{C_{p\infty}} = \frac{C_{s\infty} W_p}{M_0 - M_\infty}$$

where M_0 is the amount of migrant originally present in the polymer of weight W_p , and M_∞ is the total amount extracted at long times.

In method (2), the polymer sample is immersed in about 10 ml of extracting solvent in a typical 20 ml liquid scintillation counting vial. At specific times the sample is removed from the solvent, rinsed, and placed in another vial with fresh solvent to repeat the extraction process. The rinse is then combined with the previous extracting solvent. The total amount of migrant extracted at time t is simply the sum

of the migrant from all extracts:

$$M_t = \sum_{i=1}^t M_i$$

Method (1) is able to yield information about the equilibrium partition coefficient at an infinite extraction time. However, this method suffers from the rigid requirements of knowing accurately the weight or volume ratio of aliquot to total solution and accounting for materials lost during the sampling process for material balance purposes. As the extraction time increases, there is only a very small change in the concentration of extracted material in the solution, whereas the weighing or ratio error may persist. Therefore, the results for method (1) at long times or at high degree of extraction will show considerable scatter.

Method (2) is much simpler in operation, but simulates a condition of migration into infinite media. It is relatively free from aforementioned experimental difficulties. However, it should only be used for convenience where the migrant is highly soluble in or miscible with the solvent and thus the results of method (1) and method (2) are indistinguishable. It can neither be used to generate equilibrium partition information nor migration kinetics for cases where the migrant is sparingly soluble in the solvent.

A method often used in the literature consists of replenishing the amount of solution aliquot removed for testing with the same amount of fresh solvent, to keep the volume of solvent and exposed surface area ratio the same throughout the experiment. This method is a hybrid of method (1) and method (2), and thus suffers from the same drawbacks as those of method (2).

For both methods (1) and (2) mentioned above, the extraction vials are shaken inside a temperature controlled aluminum block on a shaking

table at a rate of about 200 reciprocations per minute.

In order to check the mass balance and variations in the migrant concentration, radioactivity of the residual low molecular weight species remaining in the polymeric sample is monitored by dissolving the sample in toluene at high temperatures after the extraction procedure is ended. We have found that the single crystals or precipitates of polymer in the counting vial do not interfere with the counting efficiency beyond the normal uncertainty of the counting results.

Experimental Run Designation

The numerical representation of experimental run designations, PPSSTTN, are assigned as follows:

- 1) Least significant digit(N)--method of observation and repetition number.
- 2) Tens and Hundreds digits(TT)--temperature in °C.
- 3) Thousands and Ten Thousands digits(SS)--solvent code.
- 4) Hundred Thousands and Millions digits(PP)--sample code.

This coding scheme is described in more detail in Table II-4 and II-5.

Estimation of Diffusion Coefficient

One of the widely used solutions for the Fickian diffusion equations solved for the case of diffusion between a plane sheet p of thickness 2ℓ and a stirred liquid s of finite volume V_s , is presented by J. Crank^[7]

$$\frac{M_t}{M_\infty} = 1 - \sum_{n=1}^{\infty} \frac{2\alpha(1+\alpha)}{1+\alpha + \alpha^2 q_n^2} \exp(-q_n^2 T) \quad (1)$$

The solution for the non-zero positive roots, q_n , of

$$\tan q_n = -\alpha q_n$$

lies between $n\pi$ when $\alpha=0$ and $(n-1/2)\pi$ when $\alpha=\infty$. At $\alpha \ll 1$,

$$q_n \sim n/(1+\alpha).$$

For other values of α ,

$$q_n \sim [n - \alpha/2(1+\alpha)]\pi$$

Table III-1.1 Migration of n-Octadecane from Linear Polyethylene

Polymer Migrant	Solvent	RUN #	WT. g.	L	cm ²	Mf, g.poly	No	Mf, solv.	DN ₀ X	CN ₂ /S
LPE .01%C18	CO	310300	0. 141	0. 072	5. 1	1. 7	0. 35	4. 4E-012	4. 1E-002	0. 1E-015
LPE .01%C18	CO	310600	0. 172	0. 069	5. 5	1. 1	0. 24	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	EN	320600	0. 157	0. 071	5. 5	1. 1	0. 22	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	EN	321600	0. 204	0. 070	5. 5	1. 1	0. 20	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	EN	323600	0. 200	0. 070	5. 5	1. 1	0. 20	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	EN	325600	0. 122	0. 071	5. 5	1. 1	0. 20	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	EN	325602	0. 122	0. 071	5. 5	1. 1	0. 20	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	HP	330241	0. 160	0. 070	5. 5	1. 1	0. 20	4. 1E-002	4. 1E-002	0. 1E-010
LPE .01%C18	HP	330301	0. 141	0. 069	4. 8	0. 91	0. 98	3. 5E-002	3. 5E-002	0. 1E-010
LPE .01%C18	HP	330601	0. 139	0. 069	4. 8	0. 91	0. 98	3. 5E-002	3. 5E-002	0. 1E-010
LPE .01%C18	OD	340301	0. 182	0. 069	6. 5	1. 3	0. 38	1. 3E-002	1. 3E-002	0. 1E-013
LPE .01%C18	OD	340601	0. 157	0. 069	6. 5	1. 3	0. 38	1. 3E-002	1. 3E-002	0. 1E-013
LPE .01%C18	TG	370300	0. 148	0. 070	5. 5	1. 6	0. 12	1. 2E-002	1. 2E-002	0. 1E-013
LPE .01%C18	TG	370600	0. 160	0. 071	5. 5	1. 6	0. 12	1. 2E-002	1. 2E-002	0. 1E-013
LPE .01%C18	AQ	380600	0. 124	0. 070	4. 2	1. 6	0. 12	1. 2E-002	1. 2E-002	0. 1E-013
LPE .01%C18	HP	430241	0. 054	0. 023	0. 0	1. 0	0. 94	1. 9E-009		
LPE .01%C18	EN	1120600	0. 331	0. 268	2. 2	1. 0	0. 90	0. 8	4. 2E-009	
LPE .01%C18	HP	1130301	0. 342	0. 258	3. 5	1. 1	0. 92	2. 9	3. 5E-009	

Table III-1.1 (Continued)

Polymer	Migrant	Solvent	RUN #	WT. g	L cm	H cm	min/cm ²	A cm ²	N ₄	solv.	D _{max} cm ² /s
LPE	C18	CO	110300	1.166	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	EN	110600	1.119	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	.5EN	1120600	1.152	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	.5EN	1125600	1.151	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	.7EN	1127600	1.151	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	.9EN	1129600	1.234	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	HP	1130240	1.449	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	HP	1130241	1.65	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	HP	1130301	1.149	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	HP	1130601	1.158	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	OD	1140301	1.158	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	OD	1140601	1.125	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	TG	1170300	1.125	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010
LPE	C18	TG	1170600	1.125	0.075	0.071	1.3E-010	1.3E-010	0.82	1.3E-010	1.3E-010

Table III-1.1 (Continued)

Polymer Migrant	Solvent	RUN #	WT. g.	A CH ₂	MnCl/ g. poly	NH ₃ / Mo	NH ₃ / solv.	DNaX CH ₂ /s
LPE	LPE	220240	0.075	2.0	2.0	2.0	2.0	2.0
1%C18	LPE	220241	0.0759	2.0	2.0	2.0	2.0	2.0
1%C18	EN	220248	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	220300	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	220601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	220602	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2225601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2225602	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2225603	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2225604	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2225605	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	229601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	2230241	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	230301	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	240601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	EN	250300	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	ON	250600	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	220248	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	220241	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	220300	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	220601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	220602	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2225601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2225602	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2225603	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2225604	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2225605	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	229601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	2230241	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	230301	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	240601	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	HP	250300	0.0760	2.0	2.0	2.0	2.0	2.0
1%C18	ON	250600	0.0760	2.0	2.0	2.0	2.0	2.0

Table III-1.1 (Continued)

Table III-1.2 Migration of n-Octadecane to/from Linear Polyethylene

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	MnCl/ g.poly	Mf/ no.	solv. %	Dmax cm ² /s	Dmin cm ² /s
LPE	0%C18 .5EN	1425600	0.223	0.064	8.0	2.4	0.07	0.0	1.6E-010	
LPE	0%C18 .5EN	1425609	0.223	0.064	8.0	0.0	0.82	0.0	4.0E-008	

Table III-1.3 Migration of n-Octadecane from Branched Polyethylene

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	MIC/ g. poly	H ₀ mol.	D _{max} cm ² /s
1%C18	BPE	710391	0.192	0.064	7.3	0.022	1.1	7E-009
1%C18	BPE	710601	0.209	0.067	7.4	0.021	1.5	4E-008
1%C18	BPE	720301	0.190	0.062	7.3	0.018	0.9	5E-008
1%C18	BPE	720601	0.190	0.064	7.0	0.019	1.0	5E-008
1%C18	BPE	730301	0.213	0.062	7.8	0.023	1.4	5E-008
1%C18	BPE	730601	0.241	0.066	7.8	0.022	1.4	5E-008
1%C18	BPE	770301	0.191	0.066	7.3	0.023	1.0	5E-008
1%C18	BPE	770601	0.171	0.064	7.4	0.022	1.0	5E-008
BPE	BPE	920399	0.131	0.057	8.6	0.022	1.0	6E-008
BPE	BPE	920600	0.162	0.054	8.6	0.021	0.9	6E-008
BPE	BPE	925300	0.154	0.055	8.6	0.021	0.9	6E-008
BPE	BPE	940301	0.155	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	940601	0.152	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	950300	0.149	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	950600	0.183	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	960300	0.094	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	960600	0.149	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	980400	0.167	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	980600	0.136	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	990300	0.191	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	990500	0.030	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	995300	0.025	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	995600	0.025	0.056	8.6	0.021	0.9	6E-008
BPE	BPE	95EN	-	-	-	-	-	-
BPE	BPE	.95EN	-	-	-	-	-	-

Table III-1.3 (Continued)

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	MIC/ g.poly	Nf/ No	% soln.	D _{max} cm ² /s
BPE	10%C18	HP	830301	0.114	0.066	4.1	0.23	9.3	1.8E-007
BPE	10%C18	OD	840301	0.098	0.065	3.6	0.24	8.3	3.2E-008
BPE	10%C18	OD	840302	0.127	0.066	4.5	0.22	10.2	2.8E-008
BPE	10%C18	OD	840601	0.099	0.065	3.7	0.23	10.0	2.2E-007

Table III-1.4 Migration of n-Octadecane to/from Branched Polyethylene

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	MICL %	NH ₃ vol v.	Dmax cm ² /s
BPE 0%C18	.SEN	1325600	0.204	0.064	7.5	2.7	0.06	1.3E-010
BPE 0%C18	.SEN	1325609	0.204	0.064	7.5	0.0	0.86	0.2 3.3E-008

Table III-1.5 Migration of n-Octadecane from Polypropylene

Polymer	Migrant	Solvent	RUN #	WT.	L	A	MICL / g. poly	NFC solv.	CN2/s	Dmax
PP	1%C18	CO	2810300	0.049	0.027	4.3	5.2	1.00	1.00	7.0
PP	1%C18	CO	2810600	0.048	0.027	3.8	5.5	1.00	1.00	6.7
PP	1%C18	EN	2820300	0.038	0.024	3.7	4.5	1.00	1.00	7.0
PP	1%C18	HP	2820600	0.049	0.027	4.2	5.4	1.00	1.00	6.7
PP	1%C18	EN	2830301	0.036	0.024	3.7	4.5	1.00	1.00	7.0
PP	1%C18	HP	2830601	0.051	0.026	4.2	5.4	1.00	1.00	6.7
PP	1%C18	HP	2860300	0.045	0.027	3.7	4.4	1.00	1.00	7.0
PP	1%C18	TB	2860600	0.047	0.027	4.2	5.4	1.00	1.00	6.7
PP	1%C18	TB	2880300	0.049	0.028	4.2	5.4	1.00	1.00	7.0
PP	1%C18	AQ	2880600	0.047	0.027	4.2	5.4	1.00	1.00	6.7
PP	1%C18	AQ	3040301	0.061	0.028	5.2	6.1	1.00	1.00	7.0
PP	10%C18	OD	3040601	0.071	0.027	6.1	6.8E-003	1.00	1.00	6.7

Table III-2.1 Migration of n-Dotriacontane from Linear Polyethylene

Polymer	Migrant	Solvent	RUN #	WT. g	L cm	H cm	Mf. %	Solv. No.	Dmax CN2/S
LPE	1%C32	CO	1910300	0.185	0.059	7.1	1.0	1.3E-012	
LPE	1%C32	CO	1910600	0.193	0.059	7.1	1.0	1.3E-012	
LPE	1%C32	EN	1920300	0.153	0.060	7.1	1.0	1.3E-012	
LPE	1%C32	EN	1920600	0.190	0.060	7.1	1.0	1.3E-012	
LPE	1%C32	.5EN	1925600	0.189	0.059	7.1	1.0	1.3E-012	
LPE	1%C32	HP	1930301	0.190	0.061	7.1	1.0	1.3E-012	
LPE	1%C32	HP	1930601	0.181	0.058	7.1	1.0	1.3E-012	
LPE	1%C32	ON	1950300	0.156	0.059	6.6	3.3	1.3E-013	
LPE	1%C32	ON	1950600	0.091	0.058	6.6	3.3	1.3E-013	
LPE	1%C32	TO	1970300	0.166	0.059	6.6	3.3	1.3E-013	
LPE	1%C32	TO	1970600	0.163	0.059	6.6	3.3	1.3E-013	
LPE	1%C32	AQ	1980600	0.059					

Table III-2.2 Migration of n-Dotriacountane from Branched Polyethylene

Table III-2.3 Migration of n-Dotriaccontane from Polypropylene

Polymer	Migrant	Solvent	RUN #	WT. g	L cm ²	H cm ²	NHCl/ g. poly	No. No.	Nf./ solv.	Dmax cm ² /s
PP	1%C32	CO	2910300	0.044	0.026	1.0	4.1	1.00	0.78	2.4E-012
PP	1%C32	CO	2910600	0.046	0.027	1.0	4.1	1.00	0.85	2.6E-011
PP	1%C32	EN	2920300	0.047	0.027	1.0	4.1	1.00	0.83	2.6E-010
PP	1%C32	EN	2920600	0.044	0.027	1.0	4.1	1.00	0.88	2.6E-009
PP	1%C32	HP	2930301	0.041	0.027	1.0	4.1	1.00	0.97	2.6E-008
PP	1%C32	HP	2930601	0.045	0.027	1.0	4.1	1.00	0.96	2.6E-007
PP	1%C32	TB	2960300	0.039	0.026	3.6	4.5	0.99	3.0E-012	
PP	1%C32	TB	2960600	0.041	0.027	3.7	4.5	0.99	3.0E-011	
PP	1%C32	AQ	2980300	0.046	0.027	4.2	4.5	0.99	3.0E-010	
PP	1%C32	AQ	2980600	0.036	0.027	3.3	4.5	0.92	3.0E-014	

Table III-3.1 Migration of BHT from Linear Polyethylene

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	NH ₃ g. poly	NH ₃ solv. No	D _{max} CH ₂ /s
LPE .01%BHT	CO	2010300	0.159	0.060	3.3	0.001	1.2E-011	1.7E-012
LPE .01%BHT	CO	2010600	0.152	0.061	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	EN	2020300	0.170	0.061	3.3	0.001	1.4E-011	1.5E-011
LPE .01%BHT	EN	2020600	0.089	0.061	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2030301	0.185	0.060	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2030601	0.192	0.061	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2030602	0.134	0.061	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	T0	2070300	0.172	0.059	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	T0	2070600	0.166	0.060	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2130601	0.058	0.025	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2210300	0.072	0.027	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	CO	2210600	0.093	0.027	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	EN	2220300	0.063	0.027	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2230300	0.075	0.027	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	HP	2230601	0.072	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	ON	2250300	0.065	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	ON	2250600	0.049	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	TB	2260300	0.062	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	TB	2260600	0.045	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	AG	2280300	0.060	0.026	3.3	0.001	1.4E-010	1.5E-010
LPE .01%BHT	AG	2280600	0.049	0.026	3.3	0.001	1.4E-010	1.5E-010

Table III-3.1 (Continued)

Polymer	Solvent	RUN #	WT. g	L cm	A cm ²	MrCi/g poly	No.	% solv.	Dmax	CN2/s
LPE	.01%BHT	CO	2610600	0.056	0.027	5.2	5.5	0.97	3.7	5.5E-010
LPE	.01%BHT	EN	2620600	0.055	0.024	5.2	5.2	0.96	0.6	6.8E-010
LPE	.01%BHT	HP	2630301	0.073	0.025	6.6	5.7	0.97	6.8	2.0E-008
LPE	.01%BHT	HP	2630601	0.053	0.025	5.2	5.6	0.97	7.0	1.1E-007

Table III-3.2 Migration of BHT from Branched Polyethylene

Polymer	Migrant	Solvent	RUN #	WT. g	L cm ²	A cm ²	MPCi/ g. poly	Mf/ Mo	% solv.	Dmax cm ² /s
BPE	.01% BHT	CO	2310300	0.071	0.062	2.9	3.0	0.70	0.6	1.7E-010
BPE	.01% BHT	CO	2310600	0.091	0.062	4.1	5.4	0.80	1.0	1.5.7E-009
BPE	.01% BHT	EH	2320300	0.155	0.060	7.1	8.0	0.82	0.8	2.2E-009
BPE	.01% BHT	EN	2320600	0.079	0.061	3.1	3.0	0.83	0.8	1.6E-009
BPE	.01% BHT	EN	2320602	0.095	0.062	3.2	3.0	0.83	0.8	1.4E-009
BPE	.01% BHT	HP	2330301	0.077	0.062	4.1	4.1	0.84	0.8	2.2E-008
BPE	.01% BHT	HP	2330601	0.081	0.062	3.3	3.5	0.78	0.7	1.7E-008
BPE	.01% BHT	HP	2330602	0.089	0.063	3.5	3.5	0.78	0.7	1.5E-008
BPE	.01% BHT	HP	2330603	0.069	0.061	2.8	2.4	0.72	0.7	1.2E-007
BPE	.01% BHT	HP	2330603	0.069	0.061	2.2	2.2	0.73	0.7	1.0E-007
BPE	.01% BHT	TO	2370300	0.070	0.061	3.1	3.0	0.76	0.8	2.0E-010
BPE	.01% BHT	TO	2370600	0.077	0.062	3.2	3.0	0.76	0.8	1.6E-009
BPE	.01% BHT	AQ	2380600	0.160	0.062	5.9	5.9	0.50	0.6	6.5E-011
BPE	.01% BHT	HP	2430601	0.058	0.028	5.2	1.7	0.12	13.0	2.3E-007
BPE	.01% BHT	HP	2430602	0.057	0.027	5.2	1.8	0.13	12.0	2.6E-007

Table III-3.2 (Continued)

Polymer	Migrant	Solvent	RUN #	WT. g	L cm2	A cm2	Mg/ g. poly	No.	solv.	Dmax cm2/s
BPE	.01%BHT	CO	2510300	0.064	0.027	0.027	0.000	10	1.0	9E-010
BPE	.01%BHT	CO	2510600	0.068	0.027	0.028	0.000	11	1.0	3E-010
BPE	.01%BHT	EN	2520300	0.077	0.027	0.026	0.000	12	1.0	1E-010
BPE	.01%BHT	EN	2520602	0.062	0.027	0.027	0.000	13	1.0	1E-010
BPE	.01%BHT	HP	2530301	0.072	0.027	0.026	0.000	14	1.0	1E-010
BPE	.01%BHT	HP	2530601	0.059	0.027	0.026	0.000	15	1.0	1E-010
BPE	.01%BHT	ON	2550300	0.066	0.038	0.026	0.000	16	1.0	1E-010
BPE	.01%BHT	ON	2550600	0.038	0.027	0.026	0.000	17	1.0	1E-010
BPE	.01%BHT	TB	2550602	0.061	0.027	0.026	0.000	18	1.0	1E-010
BPE	.01%BHT	TB	2560300	0.065	0.027	0.027	0.000	19	1.0	1E-010
BPE	.01%BHT	TB	2560600	0.071	0.027	0.027	0.000	20	1.0	1E-010
BPE	.01%BHT	TB	2560602	0.072	0.027	0.027	0.000	21	1.0	1E-010
BPE	.01%BHT	HB	2580400	0.031	0.027	0.027	0.000	22	1.0	1E-010
BPE	.01%BHT	HB	2590402	0.033	0.027	0.027	0.000	23	1.0	1E-010
BPE	.01%BHT	HB	2590600	0.045	0.027	0.027	0.000	24	1.0	1E-010
BPE	.01%BHT	HB	2590602	0.019	0.027	0.027	0.000	25	1.0	1E-010
BPE	.01%BHT	HB	2595300	0.020	0.027	0.027	0.000	26	1.0	1E-010
BPE	.95EN	EN	2595300	0.019	0.027	0.027	0.000	27	1.0	1E-010
BPE	.95EN	EN	2595600	0.020	0.027	0.027	0.000	28	1.0	1E-010

Table III-3.3 Migration of BHT from Polypropylene

Polymer	Solvent	RUN #	WT. g	L cm	A cm ²	Mf/ g.poly	Mf/ % solv.	DNUX cm ² /s
PP	.01% BHT	CO	2710390	0.052	0.027	4.7	0.30	7.3E-013
PP	.01% BHT	CO	2710600	0.046	0.027	4.2	0.37	3.6E-012
PP	.01% BHT	EN	2720390	0.035	0.027	3.5	0.39	3.2E-012
PP	.01% BHT	EN	2720600	0.035	0.026	3.5	0.38	3.6E-012
PP	.01% BHT	HP	2730391	0.040	0.026	3.5	0.39	3.5E-012
PP	.01% BHT	HP	2730601	0.041	0.026	3.5	0.75	5.0E-012
PP	.01% BHT	TB	2760390	0.039	0.027	3.5	0.83	5.0E-012
PP	.01% BHT	TB	2760600	0.035	0.027	3.5	0.16	5.0E-012
PP	.01% BHT	AQ	2780390	0.042	0.027	4.3	0.60	5.0E-012
PP	.01% BHT	AQ	2780600	0.045	0.027	4.3	0.11	5.0E-012

Table III-4.1 Migration of n-Octadecane into Triglycerides

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	MmCi/ g.poly	Hf/ No	Dmax cm ² /s
LPE	1%C18	CO	110300	0.166	0.073	5.1	1.9	3E-010
LPE	1%C18	TO	170300	0.138	0.072	4.6	2.2	1.6E-010
LPE	0.01%C18	CO	310300	0.141	0.067	5.1	1.7	1.4E-012
LPE	0.01%C18	TO	370300	0.148	0.070	5.5	2.0	0.0E-012
BPE	1%C18	CO	710301	0.192	0.064	4.0	1.1	0.0E-009
BPE	1%C18	TO	770301	0.191	0.066	4.2	1.3	4.4E-010
BPE	1%C18	TB	960300	0.094	0.057	4.3	1.2	7.7E-010
PP	1%C18	CO	2810300	0.049	0.027	4.2	1.1	4.4E-011
PP	1%C18	TB	2860300	0.045	0.027	4.5	1.1	4.4E-011
BPE	1%C18	HB	990400	0.036	0.056	1.8	0.2	0.0E-009
LPE	1%C18	CO	110600	0.119	0.075	3.9	1.6	2.0E-009
LPE	1%C18	TO	170600	0.125	0.075	4.0	1.6	8E-009
LPE	0.01%C18	CO	310600	0.172	0.072	5.4	1.2	1.2E-009
LPE	0.01%C18	TO	370600	0.160	0.071	5.7	1.2	0.0E-009
BPE	1%C18	CO	710601	0.209	0.067	4.4	1.0	2E-008
BPE	1%C18	TO	770601	0.171	0.064	4.2	1.0	2E-008
BPE	1%C18	TB	960600	0.149	0.056	4.5	1.0	2E-008
BPE	1%C18	HB	990600	0.101	0.057	4.2	0.9	1E-009
PP	1%C18	CO	2810600	0.040	0.027	4.0	0.9	1E-009
PP	1%C18	TB	2860600	0.047	0.027	4.2	0.9	1E-009

Table III-4.2 Migration of n-Octadecane into Ethanol and into n-Octanol

Polymer Migrant	Solvent	RUN #	WT. g.	L CN2	A CN2 g. poly	Mg. MMCl / g. poly	Mf. % solv.	Dmax CN2/g.
LPE 1%C18	EN	220240	0.075	0.019	0.5	2.4	0.9	2.6E-010
LPE 1%C18	EN	220241	0.041	0.018	4.9	2.3	0.9	1.2E-010
LPE 1%C18	EN	220300	0.036	0.019	4.1	2.1	0.9	1.3E-010
BPE 1%C18	EN	720301	0.190	0.062	7.4	0.95	0.5	5.5E-010
BPE 1%C18	EN	920300	0.131	0.056	5.6	0.82	0.4	6.1E-010
PP 1%C18	EN	2820300	0.038	0.024	3.8	5.4	1.00	2.0E-011
LPE 1%C18	EN	120600	0.152	0.073	5.9	1.9	0.99	5.7E-009
LPE 1%C18	EN	220600	0.059	0.019	6.6	1.1	0.93	4.0E-009
LPE 1%C18	EN	220601	0.051	0.019	6.1	1.8	0.89	1.1E-009
LPE .01%C18	EN	320600	0.157	0.069	5.3	0.66	1.00	1.0E-008
BPE .1%C18	EN	720601	0.190	0.064	7.8	0.82	0.99	1.9E-008
BPE .1%C18	EN	920600	0.162	0.057	6.0	0.27	0.99	3.0E-008
LPE .01%C18	EN	1120600	0.331	0.268	2.3	1.6	0.98	4.2E-009
PP .1%C18	EN	2820600	0.649	0.027	4.3	4.8	1.00	5.0E-009
LPE 1%C18	ON	250300	0.013	0.019	1.7	1.4	0.92	2.2E-010
BPE 1%C18	ON	950300	0.149	0.056	6.3	0.2	0.95	6.5E-010
LPE 1%C18	ON	250600	0.008	0.019	1.1	1.5	0.98	7.0E-009
BPE 1%C18	ON	950600	0.103	0.056	4.4	0.2	0.99	7.6E-009

Table III-4.3 Migration of n-Octadecane into ethanol/water Mixture

Polymer	Migrant	Solvent	RUN #	WT. g.	L cm	A cm ²	MnCl/g. poly	Mf/g. MnCl	Dmax cm ² /s
LPE .01%C18	.1EN		321600	0.204	0.071	6.7	0.7	0.00	3.7E-016
LPE .01%C18	.3EN		323600	0.200	0.070	6.7	1.1	0.00	3.0E-015
BPE 1%C18	.5EN		925300	0.154	0.057	6.5	0.2	0.02	3. 4E-012
LPE 1%C18	.5EN		125600	0.151	0.071	6.7	0.7	0.00	3.0E-019
LPE 1%C18	.5EN		125602	0.151	0.071	6.7	0.7	0.00	3.0E-019
LPE 1%C18	.5EN		125603	0.151	0.071	6.7	0.7	0.00	3.0E-019
LPE 1%C18	.5EN		225600	0.054	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		225601	0.054	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		225602	0.061	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		225603	0.061	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		225604	0.061	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		225605	0.061	0.028	6.1	0.19	0.1	6.4E-017
LPE 1%C18	.5EN		325600	0.122	0.122	6.1	0.71	0.3	1.8E-016
LPE 1%C18	.5EN		325602	0.122	0.122	6.1	0.71	0.3	1.8E-016
LPE 1%C18	.5EN		325603	0.122	0.122	6.1	0.71	0.3	1.8E-016
LPE 1%C18	.5EN		325604	0.122	0.122	6.1	0.71	0.3	1.8E-016
LPE 1%C18	.5EN		325605	0.122	0.122	6.1	0.71	0.3	1.8E-016
LPE 0%C18	.01EN		925600	0.154	0.056	6.7	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		925602	0.154	0.056	6.7	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		925603	0.154	0.056	6.7	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		925604	0.154	0.056	6.7	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		925605	0.154	0.056	6.7	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1325600	0.204	0.064	7.0	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1325602	0.204	0.064	7.0	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1325603	0.204	0.064	7.0	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1325604	0.204	0.064	7.0	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1325605	0.204	0.064	7.0	0.64	0.87	4.1E-009
LPE 0%C18	.01EN		1425600	0.223	0.064	8.0	0.64	0.82	0.58
LPE 0%C18	.01EN		1425602	0.223	0.064	8.0	0.64	0.82	0.58
LPE 0%C18	.01EN		1425603	0.223	0.064	8.0	0.64	0.82	0.58
LPE 0%C18	.01EN		1425604	0.223	0.064	8.0	0.64	0.82	0.58
LPE 0%C18	.01EN		1425605	0.223	0.064	8.0	0.64	0.82	0.58
LPE .01%C18	.7EN		127600	0.236	0.074	7.4	1.9	2.1	4.2E-009
LPE .01%C18	.9EN		129600	0.234	0.076	7.4	1.9	2.1	4.2E-009
LPE .01%C18	.9EN		229601	0.054	0.019	6.9	0.98	0.98	7.7E-009
BPE 1%C18	.95EN		995300	0.030	0.057	1.4	0.2	0.81	4.1E-010
BPE 1%C18	.95EN		995600	0.025	0.057	1.4	0.2	0.81	4.1E-010

Table III-4.5 Migration of n-Octadecane into n-Heptane

Polymer Migrant	Solvent	RUN #	WT. g.	L cm	A cm ²	MMOI/g. poly	NF% solv.	Dmax cm ² /s
LPE 1%C18	HP	130240	0.449	0.068	14.0	0.98	1.3E-008	1.1.6E-008
LPE 1%C18	HP	130241	0.165	0.070	5.8	0.98	1.2E-008	1.1.6E-008
LPE 1%C18	HP	230241	0.043	0.019	5.2	0.98	1.2E-008	1.1.6E-008
LPE .01%C18	HP	330241	0.160	0.070	5.2	0.98	1.2E-008	1.1.6E-008
LPE .01%C18	HP	430241	0.054	0.023	0.0	0.98	1.2E-008	1.1.6E-008
LPE 1%C18	HP	130301	0.149	0.071	4.8	0.98	1.3E-008	1.1.6E-008
LPE 1%C18	HP	230301	0.034	0.019	4.0	0.98	1.2E-008	1.1.6E-008
LPE 1%C18	HP	330301	0.141	0.069	4.8	0.98	1.3E-008	1.1.6E-008
LPE 5%C18	HP	530301	0.305	0.084	8.2	0.98	1.3E-008	1.1.6E-008
BPE 1%C18	HP	730301	0.213	0.062	8.8	0.98	1.3E-008	1.1.6E-008
BPE 10%C18	HP	830301	0.114	0.066	4.15	0.98	1.3E-008	1.1.6E-008
LPE .01%C18	HP	1130301	0.342	0.258	3.5	0.98	1.3E-008	1.1.6E-008
PP 1%C18	HP	2830301	0.036	0.024	3.7	0.98	1.3E-008	1.1.6E-008
LPE 1%C18	HP	130601	0.138	0.067	4.8	0.98	1.3E-008	1.1.6E-008
LPE 1%C18	HP	130602	0.150	0.069	5.8	0.98	1.3E-008	1.1.6E-008
LPE .01%C18	HP	330601	0.139	0.069	4.7	0.98	1.3E-008	1.1.6E-008
BPE 1%C18	HP	730601	0.241	0.066	8.4	0.98	1.3E-008	1.1.6E-008
PP 1%C18	HP	2830601	0.051	0.026	0.0	0.98	1.3E-008	1.1.6E-008

Table III-4.6 Migration of n-Octadecane into n-Octadecane

Polymer	Migrant	Solvent	RUN #	WT. g.	L cm	A cm ²	MnCl/ g. poly	Hf/ No	solv. %	Dmax cm ² /s
LPE	1%C18	0%C18	140301	0.138	0.072	4.069	7.303	1.7E-009	3.1E-009	3.1E-009
LPE	.01%C18	0%C18	340301	0.182	0.069	6.089	7.303	1.7E-009	3.1E-009	3.1E-009
LPE	.5%C18	0%C18	540301	0.310	0.027	3.027	7.303	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	640301	0.045	0.065	6.065	6.065	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	840301	0.098	0.127	6.066	6.066	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	840302	0.127	0.056	6.056	6.056	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	940301	0.155	0.028	6.056	6.056	1.7E-009	3.1E-009	3.1E-009
PP	10%C18	10%C18	3040301	0.061	0.018	5.018	5.018	1.7E-009	3.1E-009	3.1E-009
LPE	1%C18	0%C18	140601	0.083	0.076	3.076	3.076	1.7E-009	3.1E-009	3.1E-009
LPE	.01%C18	0%C18	240601	0.035	0.069	5.069	5.069	1.7E-009	3.1E-009	3.1E-009
LPE	.5%C18	0%C18	340601	0.157	0.085	5.085	5.085	1.7E-009	3.1E-009	3.1E-009
LPE	1%C18	1%C18	540601	0.325	0.027	1.027	1.027	1.7E-009	3.1E-009	3.1E-009
LPE	.01%C18	1%C18	640601	0.146	0.065	6.065	6.065	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	840601	0.099	0.152	6.056	6.056	1.7E-009	3.1E-009	3.1E-009
BPE	10%C18	10%C18	940601	0.152	0.071	6.056	6.056	1.7E-009	3.1E-009	3.1E-009
PP	10%C18	10%C18	3040601	0.071	0.027	1.027	1.027	1.7E-009	3.1E-009	3.1E-009

Table III-4.4 Migration of n-Octadecane into Water

Polymer Migrant	Solvent	RUN #	WT. g.	L CH ₂	A CH ₂	MgCir, g.poly	Mf% solv.	D _{max} cm ² /s
LPE .01%C18	AQ	380600	0.124	0.070	4.2	1.6	0.6	7.8E-013
BPE 1%C18	AQ	980600	0.167	0.057	6.7	0.2	0.5	8.2E-014
PP 1%C18	AQ	2880300	0.049	0.028	4.3	5.2	0.00	2.0E-017
PP 1%C18	AQ	2880600	0.047	0.027	4.3	5.0	0.00	1.0E-014

Table III-5.1 Migration of n-Dotriaconitrile into Triglycerides

Polymer	Migrant	Solvent	RUN #	WT. g	L cm	A mmCi/ ³ g. poly	Nf. %	Solv. No.	Dmax cm ² /s
BPE	1%C32	CO	1610300	0.093	0.028	7.6 115.0	0.6	3.4E-012	3.83.1.85.4.8
BPE	1%C32	TB	1660300	0.045	0.028	3.8 42.6	1.3	1.3E-012	3.83.1.85.4.8
BPE	1%C32	T0	1670300	0.074	0.028	6.3 108.0	0.6	1.3E-012	3.83.1.85.4.8
BPE	1%C32	CO	1910300	0.185	0.059	7.1 155.4.6	0.6	1.3E-012	3.83.1.85.4.8
LPE	1%C32	T0	1970300	0.166	0.059	6.5 154.4.6	0.6	1.3E-012	3.83.1.85.4.8
LPE	1%C32	CO	2910300	0.044	0.026	4.0 70.9.6	0.6	1.3E-012	3.83.1.85.4.8
PP	1%C32	TB	2960300	0.039	0.026	3.6	0.6	1.3E-012	3.83.1.85.4.8
PP	1%C32	TB	1690400	0.029	0.029	2.8 113.0	0.75	2.2	3.7E-011
BPE	1%C32	HB	1690400	0.029	0.029	2.8 113.0	0.75	2.2	3.7E-011
BPE	1%C32	CO	1610600	0.051	0.028	4.3 113.0	0.95	1.5	2.0E-009
BPE	1%C32	TB	1660600	0.032	0.029	2.7 116.0	0.95	1.5	2.0E-009
BPE	1%C32	T0	1670600	0.067	0.027	5.7 108.0	0.95	1.5	2.0E-009
BPE	1%C32	HB	1690600	0.019	0.028	1.7 115.5.0	0.95	1.5	2.0E-009
BPE	1%C32	CO	1910600	0.193	0.059	7.0 115.5.0	0.95	1.5	2.0E-009
LPE	1%C32	T0	1970600	0.163	0.059	6.3 14.1	0.95	1.5	2.0E-009
PP	1%C32	CO	2910600	0.046	0.027	3.7	0.95	1.5	2.0E-009
PP	1%C32	TB	2960600	0.041	0.027	3.7	0.95	1.5	2.0E-009

Table III-5.2 Migration of n-Dotriaccontane into Ethanol, Ethanol/Water Mixtures and n-Octanol

Polymer Migrant	Solvent	RUN #	WT. g.	L CN	A CN2 g. poly	Mf. %	solv. %	Dmax cm2/s
BPE 1%C32	EN	1620300	0.051	0.028	4.2	115.0	0.17	0.3
LPE 1%C32	EN	1920300	0.153	0.060	5.9	5.3	0.04	2.3E-012
PP 1%C32	EN	2920300	0.047	0.027	4.3	4.4	0.85	1.5E-012
BPE 1%C32	EN	1520600	0.187	0.064	7.1	93.0	0.76	1.4E-011
BPE 1%C32	EN	1620600	0.076	0.030	6.3	108.0	0.36	2.9E-009
LPE 1%C32	EN	1920600	0.190	0.060	7.5	4.2	0.63	2.9E-009
PP 1%C32	EN	2920600	0.044	0.027	4.0	5.2	0.98	5.8E-010
BPE 1%C32 .5EN	1625600	0.051	0.027	4.3	110.0	0.01	0.2	1.2E-012
LPE 1%C32 .5EN	1925600	0.189	0.059	7.4	5.2	0.00	-0.2	3.1E-014
BPE 1%C32 .95EN	1695300	0.017	0.028	1.7	115.0	0.14	0.1	1.5E-012
BPE 1%C32 .95EN	1695600	0.021	0.027	2.0	110.0	0.30	-0.5	1.9E-009
BPE 1%C32 ON	1650300	0.017	0.027	1.6	105.0	0.58	0.7	6.5E-012
LPE 1%C32 ON	1950300	0.156	0.059	6.0	5.0	0.17	2.0	2.5E-012
BPE 1%C32 ON	1650600	0.032	0.028	2.7	115.0	0.24	2.0	3.5E-009
LPE 1%C32 ON	1950600	0.091	0.058	3.9	4.9	0.80	1.9	4.7E-010

Table III-5.3 Migration of n-Dotriacontane into n-Heptane

Polymer	Migrant	Solvent	RUN #	WT. g.	L	A	MnCl ₂	Mf ₂	solv. %	D _{max}	CN ₂ /%
BPE	1%C32	HP	1530300	0.049	0.065	1.9	96.0	0.78	7.0	7.8E-010	
BPE	1%C32	HP	1630301	0.070	0.028	5.7	116.0	0.89	6.3	8.7E-010	
LPE	1%C32	HP	1930301	0.190	0.061	7.5	5.2	0.73	4.5	6.9E-010	
PP	1%C32	HP	2930301	0.041	0.027	4.0	5.1	1.00	5.8	1.4E-008	
BPE	1%C32	HP	1530601	0.191	0.065	7.2	93.0	0.99	13.0	1.7E-007	
BPE	1%C32	HP	1630601	0.069	0.028	5.7	119.0	0.99	11.0	1.6E-007	
LPE	1%C32	HP	1930601	0.181	0.059	7.2	5.1	0.95	5.3	2.7E-008	
PP	1%C32	HP	2930601	0.045	0.027	4.0	5.3	1.00	5.8	2.3E-007	

Table III-5.4 Migration of n-Dotriacontane into Water

Polymer Migrant	Solvent	RUN #	WT. g	L cm ²	A cm ²	MgCi/ g.poly	Mg/ ml.	D _{max} cm ² /s
EPE	1%C32	QQ	1680600	0.048	0.028	111.0	0.02	1.7E-015
LPE	1%C32	QQ	1980600	0.059	0.065	2.5	0.04	5.3E-013
PP	1%C32	QQ	2980300	0.046	0.027	4.2	0.08	1.3E-017
PP	1%C32	QQ	2980600	0.036	0.027	3.3	0.02	1.3E-014

Table III-6.1 Migration of BHT into Triglycerides

Polymer Migrant	Solvent	RUN #	WT. g	L	CN	A	MMCi/g	NH ₃	D _{NaX} CN2/g
LPE .01%BHT	CO	2810300	0.159	0.060	0.059	6.0	6.0	0.010	1.2E-011
LPE .01%BHT	TO	2870300	0.172	0.026	0.026	6.0	6.0	0.010	1.4E-011
LPE .01%BHT	TB	2210300	0.045	0.071	0.061	0.027	0.027	0.010	1.1E-012
BPE .01%BHT	CO	2310300	0.070	0.064	0.052	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TO	22370300	0.060	0.060	0.039	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TB	22510300	0.064	0.061	0.039	0.027	0.027	0.010	1.5E-012
PP .01%BHT	CO	2560300	0.052	0.031	0.033	0.027	0.027	0.010	1.5E-012
PP .01%BHT	TO	22710300	0.039	0.027	0.033	0.027	0.027	0.010	1.5E-012
PP .01%BHT	TB	2760300	0.039	0.027	0.033	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	HB	2590400	0.031	0.027	0.033	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	HB	2590402	0.031	0.027	0.033	0.027	0.027	0.010	1.5E-012
LPE .01%BHT	CO	2010600	0.152	0.060	0.060	0.027	0.027	0.010	1.2E-011
LPE .01%BHT	TO	2070600	0.166	0.027	0.027	0.026	0.026	0.010	1.4E-011
LPE .01%BHT	TB	2210600	0.093	0.027	0.027	0.027	0.027	0.010	1.4E-011
BPE .01%BHT	CO	22260600	0.070	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TO	22310600	0.091	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TB	2370600	0.077	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	CO	2510600	0.068	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TO	2560600	0.065	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TB	2590600	0.066	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	CO	2610600	0.056	0.027	0.027	0.027	0.027	0.010	1.5E-012
BPE .01%BHT	TB	2710600	0.046	0.027	0.027	0.027	0.027	0.010	1.5E-012
LPE .01%BHT	CO	2760600	0.035	0.027	0.027	0.027	0.027	0.010	1.5E-012
LPE .01%BHT	PP	2760600	0.035	0.027	0.027	0.027	0.027	0.010	1.5E-012

Table III-6.2 Migration of BHT into Ethanol, Ethanol/Water Mixtures, and n-Octanol

Polymer Migrant	Solvent	RUN #	WT. g	L	A cm ²	MgCl ₂ g. poly	Nf. No	Nf. solv.	Dmax cm ² /s
LPE .01%BHT	EN	2020300	0.170	0.061	6.7	3.5	0.19	7.3E-012	
LPE .01%BHT	EN	2220300	0.063	0.027	5.2	4.4	0.85	1.9E-011	
BPE .01%BHT	EN	2320300	0.155	0.060	7.5	3.9	0.92	1.0E-010	
BPE .01%BHT	EN	2520300	0.077	0.027	6.5	2.8	1.00	3.8E-010	
PP .01%BHT	EN	2720300	0.035	0.027	3.4	1.9	0.71	6.0E-012	
LPE .01%BHT	EN	2020600	0.089	0.061	3.7	3.0	0.59	0.6	2.6E-010
LPE .01%BHT	EN	2220600	0.067	0.027	5.8	4.5	0.95	0.6	2.6E-010
BPE .01%BHT	EN	2320600	0.079	0.061	3.4	3.9	0.79	0.1	6.0E-009
BPE .01%BHT	EN	2320602	0.095	0.062	3.2	3.2	0.69	1.1E-009	
BPE .01%BHT	EN	2520600	0.062	0.027	5.2	4.2	0.69	1.1E-008	
BPE .01%BHT	EN	2520602	0.072	0.029	6.0	5.2	1.00	1.5E-009	
BPE .01%BHT	EN	2520604	0.055	0.024	5.2	4.2	0.96	0.6	6.0E-010
LPE .01%BHT	EN	2620600	0.055	0.024	5.0	4.0	0.89	0.9	9.0E-010
LPE .01%BHT	EN	2720600	0.035	0.026	3.5	1.9	0.89	0.3	6.0E-010
BPE .01%BHT	.95EN	2595300	0.019	0.027	1.7	1.2	0.93	0.9	2.0E-010
BPE .01%BHT	.95EN	2595600	0.020	0.027	1.7	2.0	0.93	-0.3	6.0E-009
LPE .01%BHT	ON	2250300	0.065	0.026	5.5	3.7	0.92	1.8	1.0E-010
BPE .01%BHT	ON	2550300	0.059	0.026	5.2	1.8	0.98	1.1	5.2E-010
LPE .01%BHT	ON	2250600	0.040	0.026	3.5	3.7	0.94	2.2E-009	
LPE .01%BHT	ON	2250602	0.062	0.026	5.3	3.3	0.93	1.4E-009	
BPE .01%BHT	ON	2550600	0.066	0.027	5.4	3.5	0.98	1.4E-008	
BPE .01%BHT	ON	2550602	0.038	0.026	3.4	1.4	1.00	1.4E-008	

Table III-6.3 Migration of BHT into n-Heptane

Polymer	Solvent	RUN #	WT. g	L cm	A cm ²	MIC/ g. poly	Nf/ Mo	solv. %	Dmax cm ² /s
LPE	.01% BHT	2030301	0.185	0.060	0.022	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2230300	0.073	0.062	0.027	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2230301	0.072	0.065	0.027	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330301	0.077	0.065	0.027	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2530301	0.065	0.073	0.026	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2630301	0.040	0.026	0.026	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2730301	0.040	0.026	0.026	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2030601	0.192	0.134	0.058	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2030602	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2130601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2230601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330602	0.192	0.069	0.069	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330603	0.192	0.058	0.058	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2430601	0.192	0.057	0.057	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2430602	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2530601	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2630601	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
PP	.01% BHT	2730601	0.192	0.041	0.041	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2030601	0.192	0.134	0.058	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2030602	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2130601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2230601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330601	0.192	0.067	0.081	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330602	0.192	0.069	0.069	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2330603	0.192	0.058	0.058	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2430601	0.192	0.057	0.057	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2430602	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
BPE	.01% BHT	2530601	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
PP	.01% BHT	2630601	0.192	0.053	0.053	0.060	1.6E-008	6.7E-008	1.6E-008
LPE	.01% BHT	2730601	0.192	0.041	0.041	0.060	1.6E-008	6.7E-008	1.6E-008

Table III-6.4 Migration of BHT into Water

Polymer Migrant	Solvent	RUN #	WT. g.	L cm	A cm ²	MNCl/g. poly	Nf./Mo	% solv.	Dmax cm ² /s
LPE .01% BHT	AQ	2280300	0.069	0.028	5.8	4.8	0.32	-0.1	8.0E-013
BPE .01% BHT	AQ	2580300	0.071	0.027	5.7	1.9	0.55	0.0	2.0E-012
PP .01% BHT	AQ	2780300	0.042	0.027	3.8	1.7	0.16	0.1	3.0E-013
LPE .01% BHT	AQ	2280600	0.072	0.027	5.9	3.8	0.91	0.2	6.8E-011
BPE .01% BHT	AQ	2380600	0.160	0.062	5.9	2.3	0.50	0.0	5.6E-011
BPE .01% BHT	AQ	2580600	0.072	0.027	6.0	2.5	0.99	-0.3	3.6E-011
PP .01% BHT	AQ	2780600	0.045	0.027	4.3	1.8	0.60	0.1	1.0E-011

Table III-7.1 Activation Energies of Migration of n-Octadecane

Poly Migrant Solv Sample Solv	D30 cm ² /s	D60 cm ² /s	E kJ/mol	E kcal/mol
1%C18	110	1.3E-008	2413	2413
1%C18	130	1.6E-008	1111	1111
1%C18	140	1.6E-008	4885	4885
1%C18	170	1.8E-010	2220	2220
1%C18	220	2.2E-010	310	310
1%C18	250	2.5E-010	3340	3340
1%C18	310	3.0E-010	370	370
1%C18	340	3.4E-010	540	540
1%C18	540	5.4E-010	640	640
1%C18	710	7.1E-010	710	710
1%C18	720	7.2E-010	720	720
1%C18	730	7.3E-010	730	730
1%C18	770	7.7E-010	770	770
1%C18	840	8.4E-010	840	840
1%C18	920	9.2E-010	920	920
1%C18	940	9.4E-010	940	940
1%C18	950	9.5E-010	950	950
1%C18	960	9.6E-010	960	960
1%C18	444	4.4E-010	444	444
1%C18	2295	2.2E-011	2295	2295
1%C18	441	4.4E-011	441	441
1%C18	1112	1.1E-009	1112	1112
1%C18	1113	1.1E-009	1113	1113
1%C18	1122	1.1E-009	1122	1122
1%C18	1156	1.1E-009	1156	1156
1%C18	139	1.39E-008	139	139
LPE	110	1.1E-007	2413	2413
LPE	130	1.3E-007	1111	1111
LPE	140	1.4E-007	4885	4885
LPE	170	1.7E-007	2220	2220
LPE	220	2.2E-007	310	310
LPE	250	2.5E-007	3340	3340
LPE	310	3.1E-007	370	370
LPE	340	3.4E-007	540	540
LPE	540	5.4E-007	640	640
LPE	710	7.1E-007	710	710
LPE	720	7.2E-007	720	720
LPE	730	7.3E-007	730	730
LPE	770	7.7E-007	770	770
LPE	840	8.4E-007	840	840
LPE	920	9.2E-007	920	920
LPE	940	9.4E-007	940	940
LPE	950	9.5E-007	950	950
LPE	960	9.6E-007	960	960
LPE	444	4.4E-008	444	444
LPE	2295	2.2E-008	2295	2295
LPE	441	4.4E-008	441	441
LPE	1112	1.1E-007	1112	1112
LPE	1113	1.1E-007	1113	1113
LPE	1122	1.1E-007	1122	1122
LPE	1156	1.1E-007	1156	1156
LPE	139	1.39E-006	139	139
BPE	110	1.1E-007	2413	2413
BPE	130	1.3E-007	1111	1111
BPE	140	1.4E-007	4885	4885
BPE	170	1.7E-007	2220	2220
BPE	220	2.2E-007	310	310
BPE	250	2.5E-007	3340	3340
BPE	310	3.1E-007	370	370
BPE	340	3.4E-007	540	540
BPE	540	5.4E-007	640	640
BPE	710	7.1E-007	710	710
BPE	720	7.2E-007	720	720
BPE	730	7.3E-007	730	730
BPE	770	7.7E-007	770	770
BPE	840	8.4E-007	840	840
BPE	920	9.2E-007	920	920
BPE	940	9.4E-007	940	940
BPE	950	9.5E-007	950	950
BPE	960	9.6E-007	960	960
BPE	444	4.4E-008	444	444
BPE	2295	2.2E-008	2295	2295
BPE	441	4.4E-008	441	441
BPE	1112	1.1E-007	1112	1112
BPE	1113	1.1E-007	1113	1113
BPE	1122	1.1E-007	1122	1122
BPE	1156	1.1E-007	1156	1156
BPE	139	1.39E-006	139	139
PP	110	1.1E-007	2413	2413
PP	130	1.3E-007	1111	1111
PP	140	1.4E-007	4885	4885
PP	170	1.7E-007	2220	2220
PP	220	2.2E-007	310	310
PP	250	2.5E-007	3340	3340
PP	310	3.1E-007	370	370
PP	340	3.4E-007	540	540
PP	540	5.4E-007	640	640
PP	710	7.1E-007	710	710
PP	720	7.2E-007	720	720
PP	730	7.3E-007	730	730
PP	770	7.7E-007	770	770
PP	840	8.4E-007	840	840
PP	920	9.2E-007	920	920
PP	940	9.4E-007	940	940
PP	950	9.5E-007	950	950
PP	960	9.6E-007	960	960
PP	444	4.4E-008	444	444
PP	2295	2.2E-008	2295	2295
PP	441	4.4E-008	441	441
PP	1112	1.1E-007	1112	1112
PP	1113	1.1E-007	1113	1113
PP	1122	1.1E-007	1122	1122
PP	1156	1.1E-007	1156	1156
PP	139	1.39E-006	139	139

Table III-7.2 Activation Energies of Migration of n-Dotriacontane

Poly	Migrant	Solv	Sample	D_{30} cm ² /s	D_{60} cm ² /s	E kJ/mol	E kcal/mol
BPE	1/2C32	CO	1530	7.8E-010	1.7E-007	151	36
BPE	1/2C32	EN	1610	3.4E-012	2.2E-009	178	43
BPE	1/2C32	HP	1620	2.3E-012	1.9E-009	200	45
BPE	1/2C32	ON	1630	8.7E-010	1.6E-007	146	42
BPE	1/2C32	TB	1650	6.8E-012	1.5E-009	176	43
BPE	1/2C32	TO	1660	3.1E-012	2.1E-009	155	42
BPE	1/2C32	.95EN	1670	1.9E-012	1.9E-009	186	42
BPE	1/2C32	CO	1695	1.1E-012	1.3E-009	200	42
BPE	1/2C32	EN	1910	1.5E-012	1.9E-009	155	42
BPE	1/2C32	HP	1920	1.9E-012	2.7E-009	167	44
BPE	1/2C32	ON	1930	6.6E-010	2.4E-009	103	39
BPE	1/2C32	TB	1950	2.2E-010	3.3E-009	144	39
BPE	1/2C32	TO	1970	2.2E-010	3.3E-009	156	43
BPE	1/2C32	.95EN	2910	1.4E-011	4.3E-009	143	43
BPE	1/2C32	CO	2920	1.4E-011	3.3E-009	178	43
BPE	1/2C32	EN	2930	1.9E-012	2.3E-009	137	39
BPE	1/2C32	HP	2960	1.9E-012	1.3E-009	137	39
BPE	1/2C32	TB					

Table III-7.3 Activation Energies of Migration of BHT

Poly Migrant Solv	Solv	D30 cm2/s	D60 cm2/s	E kJ/mol	E kcal/mol
LPE	.01% BHT	2010	1.2E-012	254	25
LPE	.01% BHT	2030	1.6E-008	111	11
LPE	.01% BHT	2070	1.4E-011	165	165
LPE	.01% BHT	2210	1.1E-011	119	119
LPE	.01% BHT	2220	1.9E-008	111	111
LPE	.01% BHT	2230	2.2E-008	111	111
LPE	.01% BHT	2239	2.6E-008	111	111
LPE	.01% BHT	2250	1.0E-010	111	111
LPE	.01% BHT	2259	3.3E-012	111	111
LPE	.01% BHT	2269	8.0E-013	111	111
LPE	.01% BHT	2280	1.7E-010	111	111
LPE	.01% BHT	2310	2.6E-010	111	111
LPE	.01% BHT	2320	4.2E-008	111	111
LPE	.01% BHT	2330	4.4E-008	111	111
LPE	.01% BHT	2339	4.4E-008	111	111
LPE	.01% BHT	2370	2.0E-010	111	111
LPE	.01% BHT	2390	2.2E-010	111	111
LPE	.01% BHT	2510	5.1E-008	111	111
LPE	.01% BHT	2520	3.8E-010	111	111
LPE	.01% BHT	2539	5.3E-010	111	111
LPE	.01% BHT	2550	5.2E-010	111	111
LPE	.01% BHT	2560	3.4E-010	111	111
LPE	.01% BHT	2588	9.0E-012	111	111
LPE	.95EN	2595	2.0E-010	111	111
LPE	.95EN	2630	2.0E-008	111	111
LPE	.95EN	2710	7.3E-013	111	111

Table III-7.3 (Continued)

PP	.01% BHT	2720	6.2E-012	9.6E-010
PP	.01% BHT	2730	1.7E-008	1.7E-007
PP	.01% BHT	2760	5.5E-012	7.9E-010
PP	.01% BHT	2780	3.0E-013	1.0E-011
EH				
HP				
TB				
AQ				

**34
15
32
23**

**141
64
136
198**

may be used as the starting value in a reiterative numerical computation.

Equation (1) converges rather slowly, at small value of T, thus a sufficient number of terms must be used to avoid premature termination of the computation. At $T \ll 1$, approximately $3^{(-\log_{10} T)}$ terms are required to reach a reasonable precision. The following approaches may be used to simplify the computational effort depending on the conditions and ranges of applications.

Simplified Computation

At small values of $\alpha (< 1)$, a master curve of the quantity $M_t/(1+\alpha)M_\infty$, which is equal to M_{pt}/M_{so} for absorption or $M_{st}/\alpha M_{po}$ for extraction, as a function of T/α^2 may be constructed from equation (1). However, by limiting the computations to $0.1 > T > 0.001$, the number of terms required may be limited to about 5 at $T \sim 0.1$ and about 45 at $T \sim 0.001$ for reasonable accuracy. The values of $M_t/(1+\alpha)M_\infty$ at small α , but at the same T/α^2 , are equal to the values calculated for large α . The results of this selective computation deviates from the more rigorous and tedious computations by about 0.001% at $T \sim 0.1$, and much less than 0.001% at lower T values.

The diffusion process may be divided into three regions of T:

Region I, $T < 0.1$. The system is far from equilibrium, $M_t/(1+\alpha)M_\infty$ is a function of T/α^2 only.

Region II, $5 > T > 0.1$. The system is approaching equilibrium and detailed computation must be carried out for different α values (only 2 to 6 terms are required for equation (1) in this region). For $\alpha \ll 1$ and $T/\alpha^2 > 10^3$, $M_t/(1+\alpha)M_\infty = 1 - \alpha/(\pi T)^{1/2}$ before reaching equilibrium.

Region III, $T > 5$. For all practical purposes, equilibrium has been reached with the deviation $\delta = 1 - M_t/M_\infty \leq 10^{-T}$, where $M_\infty = M_{so}/(1+\alpha)$ for

the case of absorption and $M_\infty = M_{po}\alpha/(1+\alpha)$ for the case of extraction.

For $\alpha < 1$, equilibrium may be reached much earlier.

Computation for Migration into Infinite Media

When $\alpha \rightarrow \infty$, M_t/M_∞ approaches a limit, and equation (1) can be reduced to a function of T only:
$$M_t/M_\infty = 1 - 2 \sum_{n=1}^{\infty} \frac{1}{q_n^2} \exp(-q_n^2 T) \quad (2)$$

where $q_n = (n-1/2)\pi$. For $T < 0.1$, M_t/M_∞ is a linear function of $T^{1/2}$.

$$M_t/M_\infty = 2(T/\pi)^{1/2} = 1.128379 T^{1/2} \quad (3)$$

Deviations of equation (3) from equation (2) is less than 10^{-10} at $T < 0.05$, about 10^{-6} at $T = 0.1$, about 5×10^{-4} at $T = 0.2$ and becomes much greater than 0.01 at $T = 0.3$ or higher.

Alternate Approximation

An alternative form of the solution^[7]

$$M_t/M_\infty = (1+\alpha) [1 - e^{T/\alpha^2} \operatorname{erfc}(T^{1/2}/\alpha)], \quad (4)$$

may be used in some cases and is relatively simple to compute. One of the rational approximations^[8] for the error function yields the following

$$M_t/(1+\alpha)M_\infty = 1 - \sum_{n=1}^5 a_n \tau^n + \varepsilon \quad (5)$$

where $\tau = 1/(1+0.3275911 T^{1/2}/\alpha)$, $a_1 = 0.254829592$, $a_2 = -0.28449636$, $a_3 = 1.421413741$, $a_4 = -1.453152027$, $a_5 = 1.061405429$, and $|\varepsilon| < 1.5 \times 10^{-7}$.

At $\alpha \ll 1$, results computed from equation (2) or (3) deviates less than 0.0001 from that of equation (1) at $T/\alpha^2 < 5$ or at $M_t/(1+\alpha)M_\infty < 0.75$. Maximum deviation of 0.0035 occurs at $T/\alpha^2 \sim 500$ or at $M_t/(1+\alpha)M_\infty \sim 0.97$.

Therefore, equation (4) or (5) may be used to generate the master curve for equation (1) at $T < 0.1$ (Region I). However equations (4) or (5) do not yield any information about the region approaching equilibrium at larger values of α , and hence should be used with discretion.

By combining Equation (4) or (5) at $T < 0.1$ (Region I) and Equation (1) at $T > 0.1$ (Region II and III), computational requirements for the solution of the diffusion equation between a plane sheet and a well stirred liquid may be reduced to a minimum.

Diffusion coefficient at any point may be estimated from the above computation method by means of regression.

Effect of Swelling

The diffusion coefficients in the tables are calculated based on the original measured thickness of the sample, which is also given in the tables. The final sample is usually swollen with solvent. For volatile solvents, the absorbed solvent will leave the polymer soon after the sample is taken out of the extracting solvent. In these cases, the final sample weight is determined immediately after the surface of sample has been wiped free of the solvent. The dimensions of the final sample are generally not measured, however they may be estimated from the amount of solvent absorption by assuming an isotropic expansion. The diffusion coefficient may be adjusted to reflect the increase in the thickness as follows:

$$\begin{aligned} D &= D_0 \left(\frac{L}{L_0} \right)^2 = D_0 \left(\frac{V}{V_0} \right)^{2/3} \\ &= D_0 \left[1 + \frac{\delta_f}{1-\delta_f} \cdot \frac{\rho_0}{\rho_s} \right]^{2/3} \end{aligned}$$

where $\delta_f = (W_f - W_0)/W_f$. D , L , V , M and ρ are the diffusion coefficient, thickness, volume, weight and density respectively. The subscripts o , f , and s denote the original polymer, final polymer and solvent respectively. The sum of fractional weight increase δ_f and the fractional decrease in weight due to the migrant loss is given also in the table and indicates the weight percent of solvent content in the final sample. Therefore, the diffusion coefficients based on the final sample thickness

are increased from those based on the original thickness by approximately the same order of magnitude as the increases in weight.

Table II-1
Characteristics of Polyolefin Samples

		Linear Polyethylene NBS-SRM 1475	Branched Polyethylene NBS-SRM 1476	Isotactic Polypropylene Pro-fax 6301*
Molecular Weights	M_n	18,310 (GPC)		
	M_w	53,070 (GPC)		290,000
	M_w	52,000 (LS)		
	M_w/M_n			11
Limiting Viscosity Number	CN	0.890	0.8132	
[η], dl/g	TCB	1.010	0.9024	
	DHN	1.180	1.042	2.1
Melt Flow Rate, g/10 min		2.07	1.19	
Density, g/cm ³		0.97844	0.9312	
Isotacticity, %				95-96

GPC -- Gel Permeation Chromatography

LS -- Light Scattering

CN -- 1-chloronaphthalene

TCB -- 1,2,4-trichlorobenzene

DHN -- decahydronaphthalene (Decalin)

t [η] = 130°C for polyethylenes, 135°C for polypropylene

Melt Index by Procedure A, ASTM Method D 1238-65T, Test Condition D, 190°C, load 325 g for SRM 1475 and 1260 g for SRM 1476.

Density by ASTM Method D 1505-67; sample prepared by Procedure A, ASTM Method D 1928-68.

Isotacticity is determined as fractional insoluble in Decalin at room temperature after the entire sample has been dissolved at 160°C and allowed to cool.

*Certain commercial materials and equipment are identified in this paper to adequately specify the experimental procedure. This identification does not imply recommendation or endorsement by the National Bureau of Standards, nor does it imply that the material or equipment identified is necessarily the best available for the purpose.

Table II-2

Characteristics of Ethylene-Vinyl Acetate Copolymers

Designation	Vinyl Acetate Content %	M_n	M_n	Melt Index g/10 min	Density g/cm ³
Petrothene NA294-00	5.1	15000	89300	1.9	0.925
Ultrathene UE657-00	13.4	18700	103000	0.46	0.935

Table II-3
Characteristics of Radioactive Tracers

	<u>$\mu\text{C}_i/\text{mg}$</u>	<u>ng/25 dpm</u>
n-C ₁₈ H ₃₈ -1- ¹⁴ C	86.0	0.13
	13.3	0.83
n-C ₃₂ H ₆₆ -16,17- ¹⁴ C	45.5	0.25
3,5-di-tert-butyl-4 hydroxytoluene-7- ¹⁴ C (BHT)	57.9	0.19

Table II-4

Solvent <u>$\times 10^3$</u>	Temperature <u>$\times 10^1$</u>	Method <u>$\times 10^0$</u>
10, Corn Oil (CO)	24, 24°C	0, Limited Solvent Volume (L)
20, Ethanol (EN)	30, 30°C	1, Unlimited Solvent Volume (U)
	40, 40°C	2-8, Duplicate (or Renewed Extractions)
21, 10% Ethanol (.1 EN)	60, 60°C	9, Absorption Experiment (A)
23, 30% Ethanol (.3 EN)		
25, 50% Ethanol (.5 EN)		
27, 70% Ethanol (.7 EN)		
29, 90% Ethanol (.9 EN)		
30, n-Heptane (HP)		
40, n-Octadecane (OD)		
50, n-Octanol (ON)		
60, Tributyrin (TB)		
70, Trioctanoin (TO)		
80, Water (AQ)		
90, HB307 (HB) Synthetic Triglyceride Mixture		
95, 95% Ethanol (.95 EN)		

Letters in parentheses are abbreviations used in tables.

Table II-5

<u>Sample Code</u>	<u>Polymer</u>	<u>Migrant</u>	<u>Nominal Migrant Conc., %</u>	<u>Sp. Act. μCi/ g. Polymer</u>	<u>Thickness cm</u>
x10 ⁵					
1	LPE	C18	0.64	2.1	0.072
2	LPE	C18	0.63	2.2	0.019
3	LPE	C18	0.013	1.6	0.070
4	LPE	C18	0.01	0.96	0.020
5	LPE	C18	5.2	0.14	0.084
6	LPE	C18	5.5	0.13	0.027
7	BPE	C18	0.85	0.020	0.064
8	BPE	C18	9.8	0.23	0.066
9	BPE	C18	0.85	0.17	0.057
11	LPE	C18	0.009	1.1	0.263
13	BPE	C18	0		0.064
14	LPE	C18	0		0.064
15	BPE	C32	1.86	93.	0.064
16	BPE	C32	2.26	113.	0.028
17	BPE	C32	0		
18	LPE	C32	0		
19	LPE	C32	1.0	5.2	0.059
20	LPE	BHT	0.0062	3.6	0.060
21	LPE	BHT	0.0069	4.0	0.025
22	LPE	BHT	0.0072	4.2	0.027
23	BPE	BHT	0.0052	3.0	0.061
24	BPE	BHT	0.0030	1.8	0.028
25	BPE	BHT	0.0045	2.6	0.027
26	LPE	BHT	0.0095	5.5	0.025
27	PP	BHT	0.0029	1.8	0.027
28	PP	C18	0.82	4.9	0.027
29	PP	C32	0.97	5.3	0.027
30	PP	C18	6.8	0.16	0.028
31	E-5% VA	BHT	0.0091	5.3	0.029
32	E-13% VA	BHT	0.0057	3.3	0.069

References

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2. Smith, L. E., Sanchez, I.C., Chang, S. S. and McCrackin, F. L., "Models for the Migration of Paraffinic Additives in Polyethylene," NBSIR 79-1598, 1979.
3. Smith, L. E., Sanchez, I. C., Chang, S. S., McCrackin, F. L., and Senich, G. A., "Models for the Migration of Additives in Polyolefins," NBSIR 79-1779, 1979.
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7. Crank, J. "The Mathematics of Diffusion," Oxford University Press, 1975.
8. Abramowitz, M. and Stegun, I. A., "Handbook of Mathematical Functions," AMS 55, National Bureau of Standards, GPO, 1964.

III. Results of Polyolefin Extraction Experiments

The results of extraction experiments are summarized in Table III.1 through III.3 for the extraction of n-octadecane, n-dotriacanthane and BHT by triglycerides, ethanol, n-octanol, ethanol/water mixtures, n-heptane, n-octadecane, and water.

The tables provide information on the test pieces as to their weight, thickness L, exposed area A, specific activity of polymer, amount extracted at the end of experiment M_f/M_0 , the amount of solvent retained in the final sample and the observed maximum of the diffusion coefficient.

The value for M_f/M_0 in the tables is the final fraction of total additives migrated at the end of the experiment. It may be either an equilibrium value or a non-equilibrium value after a long time (6 months to one year) lapse. In the case of BHT experiments, most of the M_f values for migration into good solvents denote the available fraction of BHT in the sample, which varies with the conditions underwhich the sample was prepared. Therefore, in calculating diffusion coefficients for samples containing BHT, an adjusted M_0 is assigned to each sample according to the equilibrium (M_f/M_0) values observed in the good solvents. For the case of $n-C_{32}H_{66}$ in polypropylene or other samples which exhibit surface blooming of migrants, the total amount of migrant is also adjusted for the amount of migrant on the surface, before diffusion coefficients calculating from it.

Tables III-1 through III-3 summarize the experimental results arranged by migrant or additive and then according to the type of polymer, additive concentration, sample thickness, and, finally, solvents and temperature. In order to provide easy cross-references, Tables III-4

through III-6 summarize the experimental results which are arranged first by solvent and then by additive and other parameters.

The activation energies, E, calculated by a simple Arrhenius equation:

$$D = A \exp (-E/RT)$$

are given in Table III-7.

IV. Loss of BHT from Polymers in Ambient Atmosphere

ADL, Inc. had reported through private communications concerning the high volatility of BHT from their samples. When we examined our observed total BHT contents from various experiments over a 1 1/2 year period on a branched polyethylene sample, code 25, containing 45 ppm of BHT, we found a rather noticeable trend of decreasing radioactivities as a function of starting date of the experiment, Figure IV-1, although the sample plaque was always wrapped in aluminum foil except during short periods when pieces were cut from the plaque.

The volatility of BHT is further indicated by the following experiment. A piece of E-13% VA copolymer containing 60 ppm of BHT (sample code 32), 61 mg in weight with about 2 cm^2 of surface area, was placed in a sealed 20 ml vial. The bottom of the vial was kept at 30°C and the vial cap at 22-23°C. After 63 hours it was found that the aluminum cap liner had a radioactivity of 0.51% of the test piece and the glass wall of the vial had a radioactivity of 0.95% of the test piece. Thus in a relatively short period, 1.5% of the BHT was sublimed from the test piece at rather low temperatures in a closed vessel.

A further test was performed on both sample plaques 25 and 32. Four pieces were cut from each of the sample plaques. These four pieces were subjected to the following tests:

- (A) The test piece was dissolved in o-xylene to check the current specific activity.
- (B) The test piece was extracted immediately by ethanol at 60°C for three days. The residual specific activity and the activity of the extract were recorded.
- (C) The test piece was suspended in a ventilation hood at room temperature for three days before procedure (B) was performed.

(D) The test piece was placed in a vial. Forced air was blown into the vial for three days at room temperature before procedure (B) was performed.

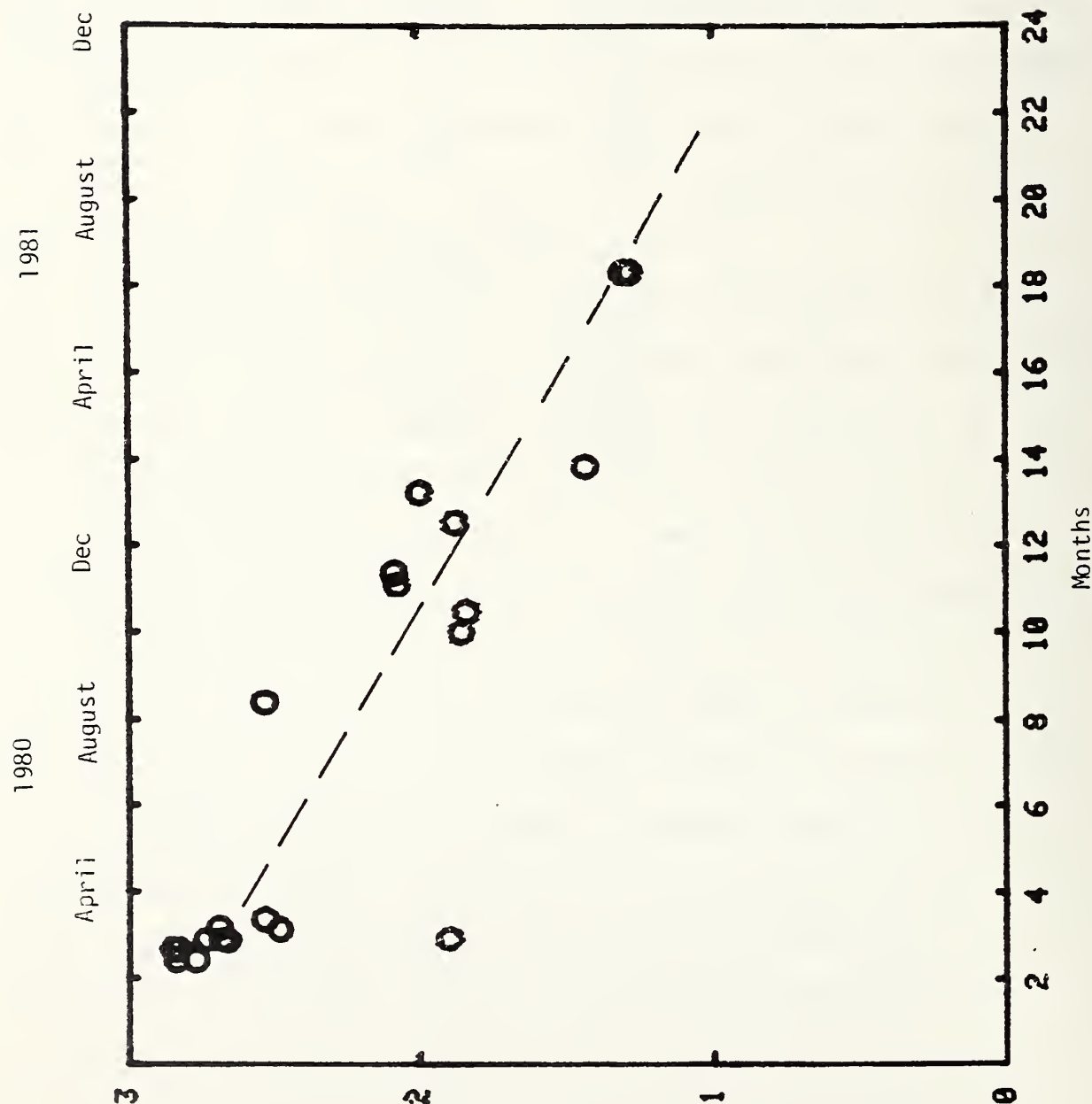
The results of these tests are presented in Table II-1. The test pieces A and B were used as controls. Apparently 30 to 50% BHT is lost from thin sheets of BPE or E-VA copolymer in mobile air at room temperature for three days. The specific activity of sample code 25, BHT in BPE, at the time of these tests was only 50% of the original specific activity of 2.8 μ Ci/g polymer, determined 16 months earlier.

These experiments and our prior experience indicate that: (1) the amount of additive remaining in the polymer after processing to produce plaques is a fraction of the additive mixed into the raw polymer, (2) for certain reactive additives, only a fraction of additive in the polymer may be extractable, the rest may be chemically bound to the polymer, (3) the total activity of the test pieces should be checked for each experiment not only to measure the uniformity of the additive concentration but also to measure the change in additive content as a function of time. Therefore reports which the results are normalized to the amount of additive supposedly added to the polymer or to the total amount of additive present in the polymer when it was manufactured, without paying further attention to the amount of additive either reacted or lost, may yield seriously incorrect kinetics of migration.

Table IV-1

Percent of Original
Activity
(Average of A + B)

Sample Code	Test Piece	Specific Activities, $\mu\text{Ci/g}$ polymer		
		Residue	Total	
25 BHT in BPE	A	--	1.28	100
	B	0.042	1.30	100
	C	0.053	.93	72
	D	0.052	.89	69
32 BHT in E-13% VA	A	--	3.29	100
	B	0.077	3.45	100
	C	0.081	2.18	65
	D	0.068	1.49	44



Specific Activity, $\mu\text{Ci/g Polymer}$

Figure IV-1

Loss of BHT from Sample 25, BHT/BPE

V. Migration of BHT from Ethylene-Vinyl Acetate Copolymers

Ethylene-vinyl acetate (E-VA) copolymers of relatively low vinyl acetate contents have been used in increasing quantities in recent years as food-contact polymer films. The migration of the antioxidant BHT from two copolymers having 5 and 13% vinyl acetate content into corn oil, anhydrous ethanol, 95% ethanol, water and n-heptane was observed with ^{14}C -labeled BHT. The characteristics of these polymers were shown earlier in Table I-2. The characteristics of the additive BHT were presented earlier in Table I-3. The sample plaques were prepared in similar manner to those of polyethylene and polypropylene samples described earlier, except that lower molding temperatures were used for the E-VA copolymers. The E-5% VA and E-13% VA samples were designated as plaque 31 and 32 with thickness of 0.027 and 0.068 cm and BHT contents of 9 and 57 PPM, respectively.

The results for individual extraction experiments are summarized in Table V-1. E-VA copolymers absorb a fair amount of solvents as shown in the table. The listed values are fractional weights of solvent contained in the final sample. Thus for experiment 3230600, the final sample contained n-heptane amounting to about 120% of the original weight of the polymer. The 12% corn oil content for sample 3210600 is higher than all other corn oil experiments.

The results of all extraction experiments for individual E-VA copolymers are shown graphically in Figures V-1.1 through V-1.4 for the migration of BHT from E-5% VA at 30 and 60°C, and from E-13% VA at 30 and 60°C, respectively. There are three illustrations for each figure, i.e., (A) M_t/M_0 is plotted against t/L^2 in a log-log plot to indicate the migration behavior over several orders of magnitudes in both M_t/M_0 and time, (B) M_t/M_0 is plotted against $t^{1/2}/L$ for the short time region

to indicate the behavior observed during fast extractions, and (C) M_t/M_0
is plotted against $t^{1/2}/L$ for the entire length of time of experiments.

From both Tables V-1 and Figures V-1, the residual BHT content in the 5% vinyl acetate copolymer samples, code 31, at the end of extraction are seen to vary from 7 to 12%. It is estimated from graphical extrapolation that approximately 7-8% of the original BHT content is not extractable even by good solvents, such as n-heptane. Almost all of the BHT content in the 13% vinyl acetate copolymer is extractable. The 5% water content in 95% ethanol has no noticeable effect on the BHT extractability for the copolymers. No partitioning of BHT was observed between the copolymers and solvents such as n-heptane, corn oil, anhydrous ethanol and 95% ethanol, due to the near-complete of extraction of all available BHT from the copolymers by these solvents.

Figures V-2.1 through 5 show the migrational behavior of BHT from E-VA copolymers into various solvents, n-heptane, corn oil, ethanol, 95% ethanol and water. These illustrations offer comparisons of behavior among different E-VA copolymers and/or at different temperatures of extraction.

In a more concise comparison of this behavior, the apparent maximum diffusion coefficients of the swollen polymers are listed in Table V-2 together with the activation energies for diffusion. In all cases except water, the diffusion coefficient for the 13% vinyl acetate copolymer is about 2 to 3 times greater than that for the 5% copolymer. Increasing in the temperature from 30 to 60°C generally increases the diffusion coefficient by 10 to 20 fold, although the changes for the solvent n-heptane are only on order of 4 to 6.

The diffusion coefficients in 95% ethanol are only slightly less than those for anhydrous ethanol, about 10-15% less for 5% vinyl acetate copolymers and about 25-45% less for 13% copolymer. Both diffusion

coefficients for experiments in anhydrous ethanol and 95% ethanol are greater than those for corn oil by similar magnitudes.

The migrational behavior of BHT from the two E-VA copolymer into water is rather unusual, as can be seen in figure V-2.5. The $\log(M_t/M_0)$ versus $\log(t/L^2)$ representation in Figure V-2.5 is changed for Figures V-3-B and V-3-C into the more familiar M_t/M_0 versus $t^{1/2}$ representation. Figure V-3-B is the short-time portion of Figure V-3-C which covered the entire time scale of experiment. These results indicate that the migration of BHT from E-VA copolymer into water occurs in two distinct stages, and that in contrast to the observations for other solvents, the diffusion coefficients of BHT from E-13% VA copolymer are slower than those from E-5% VA copolymer in the first stage. In the second stage the diffusion coefficients from E-13% VA are apparently approaching those from E-5% VA and can even surpass the latter when the time of experiment is lengthened to allow equilibrium extraction.

In the first stage, 10-50 hours, approximately 15% of the BHT from E-5% VA copolymer and about 1% of the BHT for E-13% VA copolymer, were extracted by water before the release of BHT levels off into a second stage behavior. It has not been determined whether the second stage of extraction, with its distinctly different diffusion coefficient, is due to the distribution of BHT in phase-separated regions of the copolymers or due to the reaction of water and BHT over prolonged periods.

Another plausible explanation for the two stage behavior would be surface blooming, or a very high migrant concentration near the surface layer, although the time lapse of 10 to 50 hours seems to be longer than usually needed to remove migrant from the surface or surface layer. By extending the second stage behavior to $t=0$, one may estimate from

Figure V-3-B that there might exist surface blooming of BHT amounts to about 7.5 and 11% for the E-5% VA copolymer and 0.6 and 0.57% for the E-13% VA copolymer, for 30 and 60°C experiments, respectively.

The diffusion coefficients presented in Table V-2 were calculated by first adjusting the original loading, M_0 , to the available BHT, M_0^* , when necessary. The first stage diffusion coefficients were calculated based on the above mentioned percentages for M_∞/M_0^* . By subtracting the first stage M_∞ , normalizing and then applying equilibrium partitioning when applicable, the second stage difficient coefficients were estimated.

Equilibrium partitioning of BHT between the copolymers and solvents was observed only for the case of E-5% VA and water at 60°C. In all other experiments with water, a time lapse of six months was not long enough to establish an equilibrium conditions.

Table V-1.1 Migration of BHT from E-5% VA Copolymer

Polymer	Migrant	Solvent	RUN #	WT. g.	L cm	A cm ²	NMCI/g. poly	H ₂ O solv.	DNAx CM ² /s
EVA. 05	BHT	CO	3110300	0.033	0.038	2.8	0.88	0.88	0.88
EVA. 05	BHT	CO	3110600	0.032	0.028	2.5	0.88	0.86	0.87
EVA. 05	BHT	EN	3120300	0.039	0.028	2.5	0.88	0.86	0.87
EVA. 05	BHT	EN	3120600	0.035	0.029	2.5	0.88	0.86	0.87
EVA. 05	BHT	HP	3130300	0.032	0.029	2.5	0.88	0.86	0.87
EVA. 05	BHT	HP	3130600	0.030	0.029	2.5	0.88	0.86	0.87
EVA. 05	BHT	.95EN	3195300	0.064	0.027	2.5	0.88	0.86	0.87
EVA. 05	BHT	.95EN	3195600	0.062	0.028	2.5	0.88	0.86	0.87
EVA. 05	BHT	AQ	3180300	0.036	0.030	2.6	0.88	0.86	0.87
EVA. 05	BHT	AQ	3180600	0.028	0.030	2.4	0.71	0.71	0.71

Table V-1.2 Migration of BHT from E-13% VA Copolymer

Polymer Migrant	Solvent	RUN #	WT. g	L cm	A cm ²	mmCi/g. poly.	% solv.	DNA _X cM ² /s
EVA. 13	BHT	CO	3210300	0.137	0.072	4.7	3.7	3.7E-009
EVA. 13	BHT	CO	3210600	0.121	0.067	4.5	3.2	4.2E-008
EVA. 13	BHT	EN	3220300	0.158	0.072	5.4	1.0	1.0E-008
EVA. 13	BHT	EN	3220600	0.136	0.071	4.6	1.0	1.0E-007
EVA. 13	BHT	HP	3230301	0.130	0.068	4.7	0.9	1.3E-007
EVA. 13	BHT	HP	3230601	0.127	0.070	5.0	0.9	1.3E-006
EVA. 13	BHT	.95EN	3295300	0.157	0.067	5.8	1.6	1.5E-006
EVA. 13	BHT	.95EN	3295600	0.156	0.066	5.6	1.5	1.5E-005
EVA. 13	BHT	AQ	3280300	0.147	0.068	5.5	1.2	1.2E-013
EVA. 13	BHT	AQ	3280600	0.123	0.068	4.4	3.0	3.0E-012

Table V-2

Diffusion Coefficients and Activation Energies of
Migration of BHT from Ethylene-Vinyl Acetate Copolymers into Various Solvents

Solvent	$D_{\max} (10^{-8} \text{ cm}^2 \text{s}^{-1})$				E	
	30°C		60°C		kJ mol^{-1}	
	E5%VA	E13%VA	E5%VA	E13%VA	E5%VA	E13%VA
n-Heptane	14	30	57	180	39	51
Corn Oil	0.15	0.37	2.3	4.9	76	72
Ethanol	0.19	1.0	3.7	10	83	65
95% Ethanol	0.19	0.55	3.2	7.5	82	73
Water - 1st Stage	0.0046	0.00007	0.005	0.00034	67	44
2nd Stage	0.000096	0.000023	0.0028	0.0010	94	106

- Figure V-1.1-A, B, C Migration of BHT from E-5% VA Copolymer at 30°C
Figure V-1.2-A, B, C Migration of BHT from E-5% VA Copolymer at 60°C
Figure V-1.3-A, B, C Migration of BHT from E-13% VA Copolymer at 30°C
Figure V-1.4-A, B, C Migration of BHT from E-13% VA Copolymer at 60°C

Legends for Figure V-1.1

Corn Oil	△
Ethanol	○
95% Ethanol	○
n-Heptane	+
Water	*

Figure V-1.1-A

MIGRATION OF BHT
FROM ETHYLENE-5% VINYL ACETATE COPOLYMER
AT 30°C

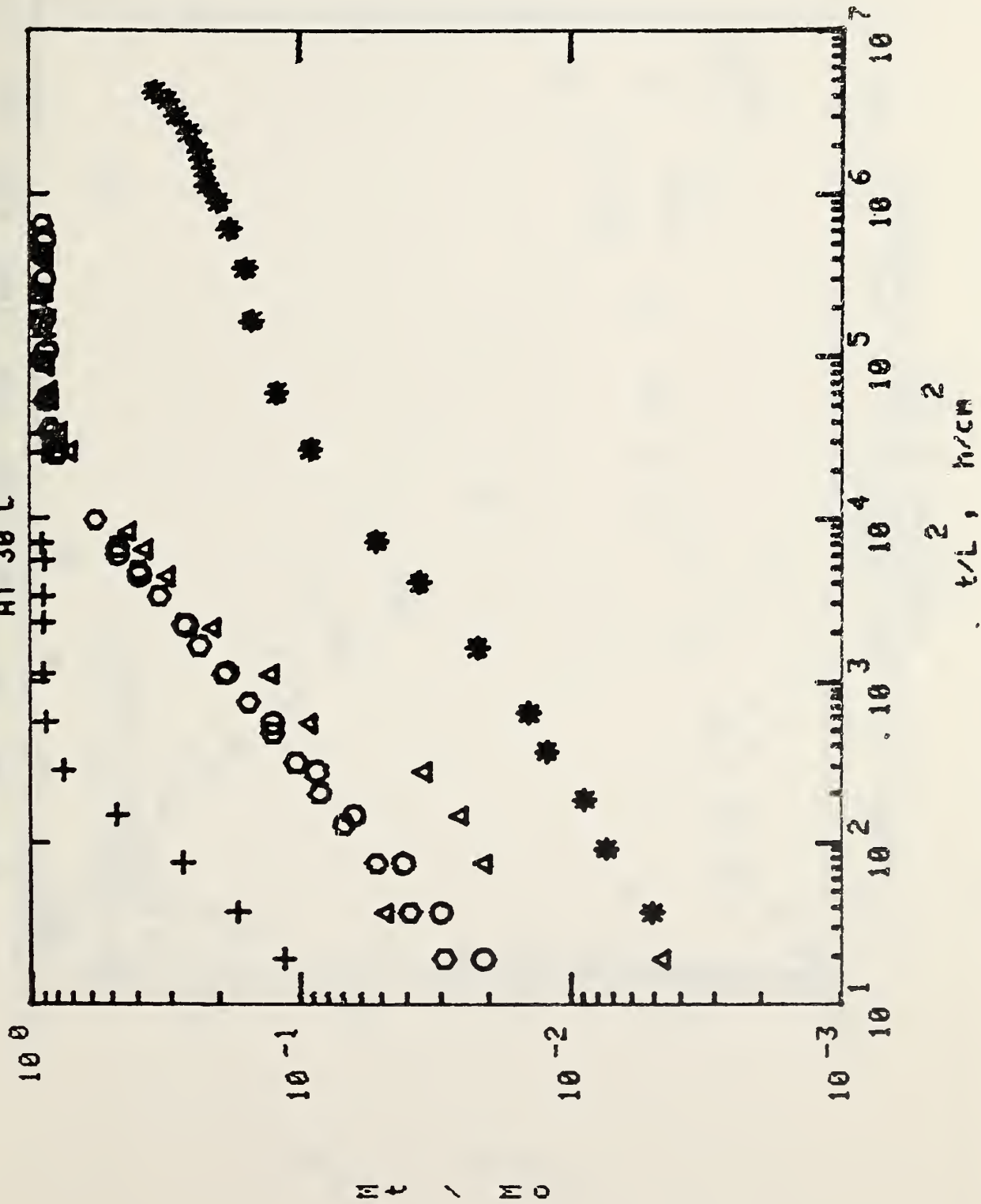


Figure V-1.1-B

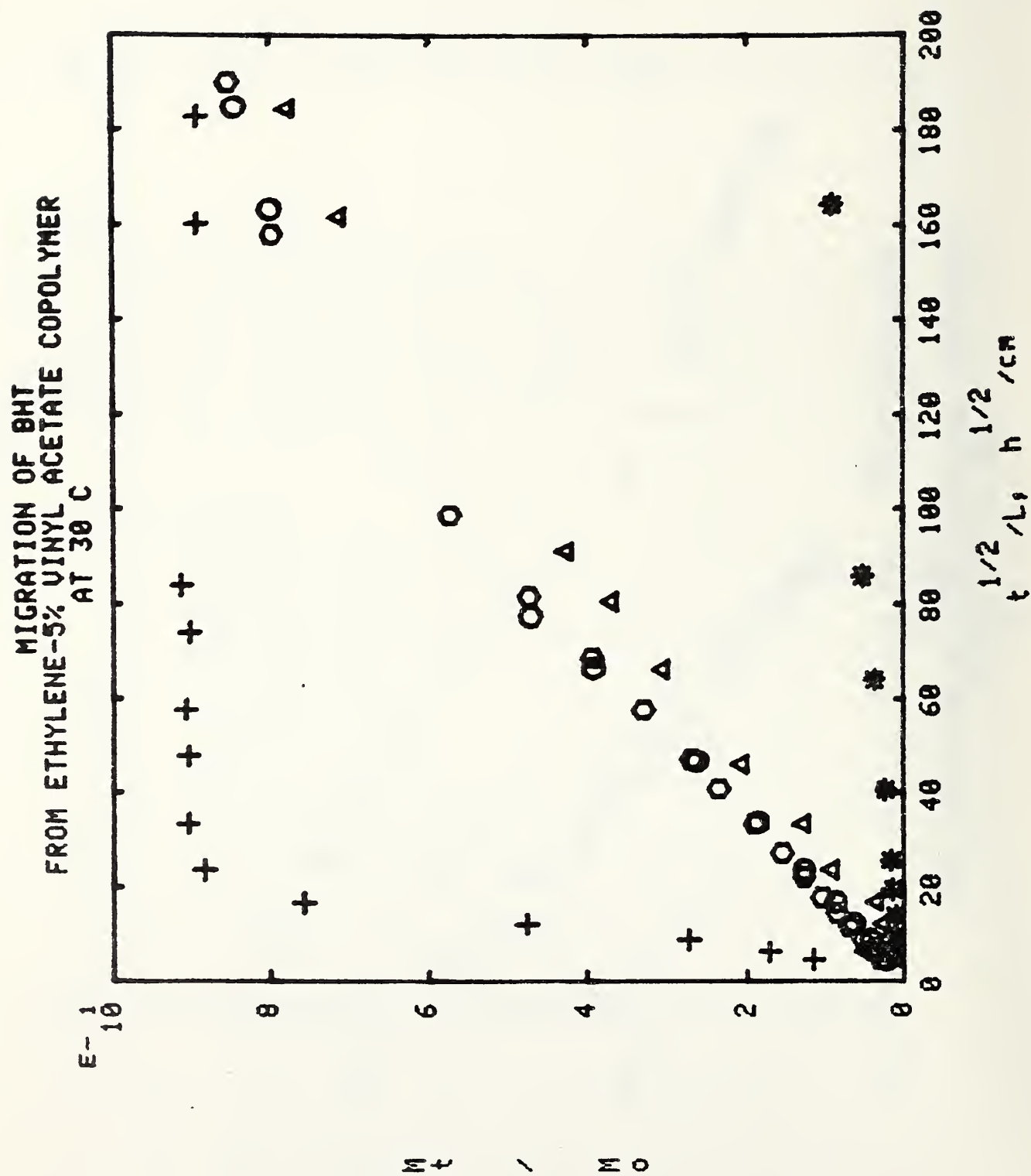


Figure V-1.1-C

FROM ETHYLENE-5% VINYL ACETATE COPOLYMER
AT 30°C

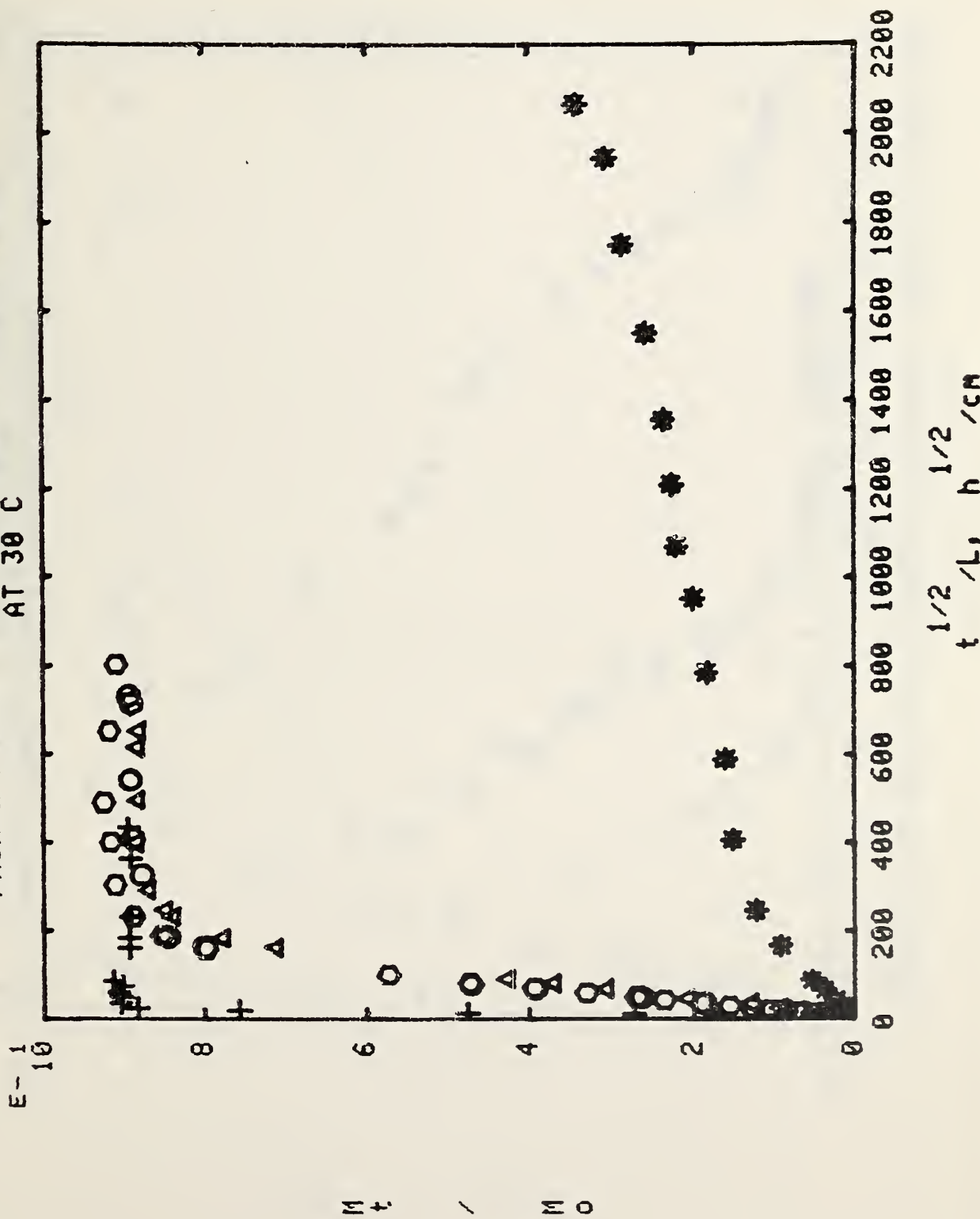


Figure V-1.2-A

MIGRATION OF SHT FROM ETHYLENE-5% VINYL ACETATE COPOLYMER
AT 60 °C

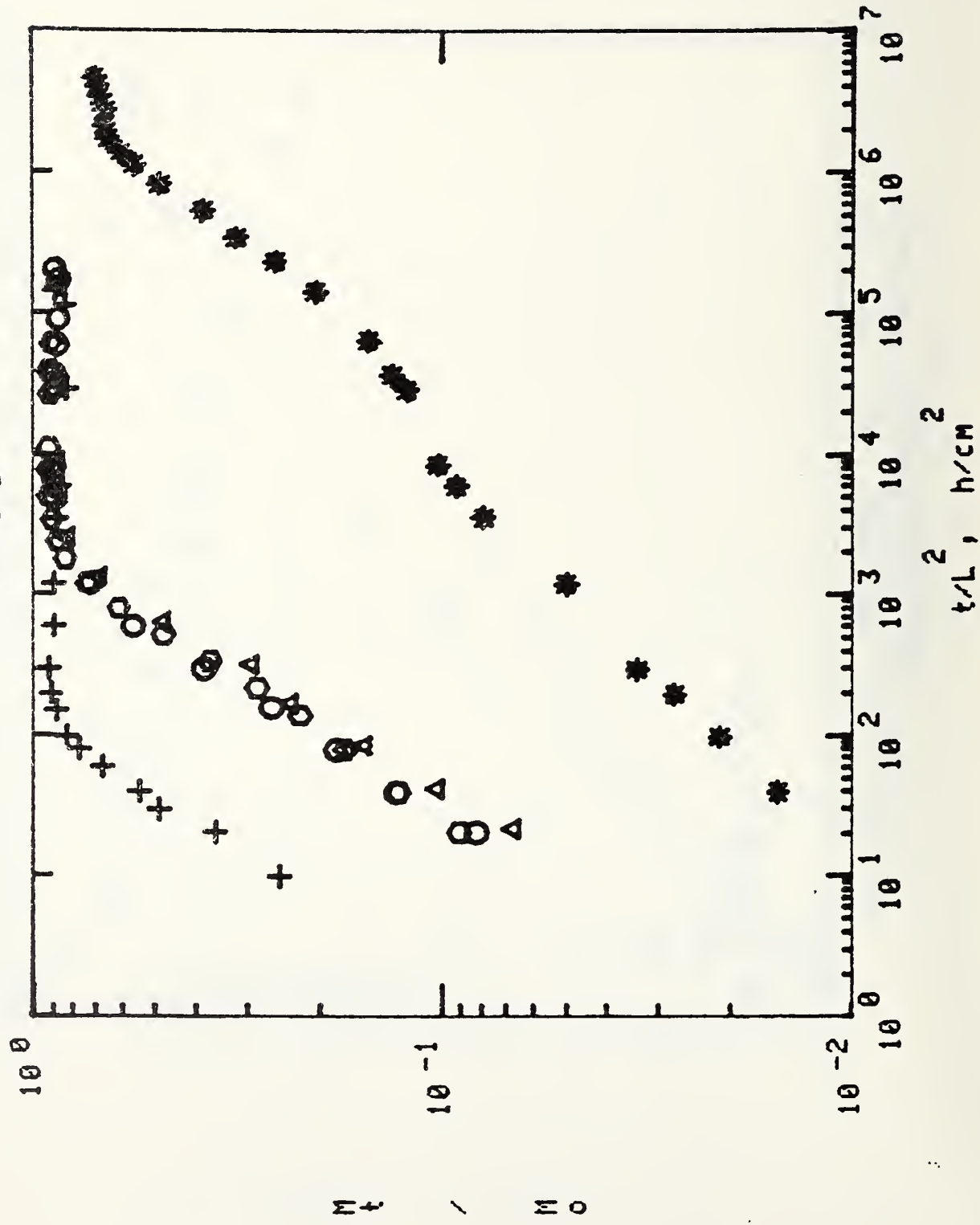


Figure V-1.2-B

MIGRATION OF BHT FROM ¹ETHYLENE-5% VINYL ACETATE COPOLYMER
AT 60 °C

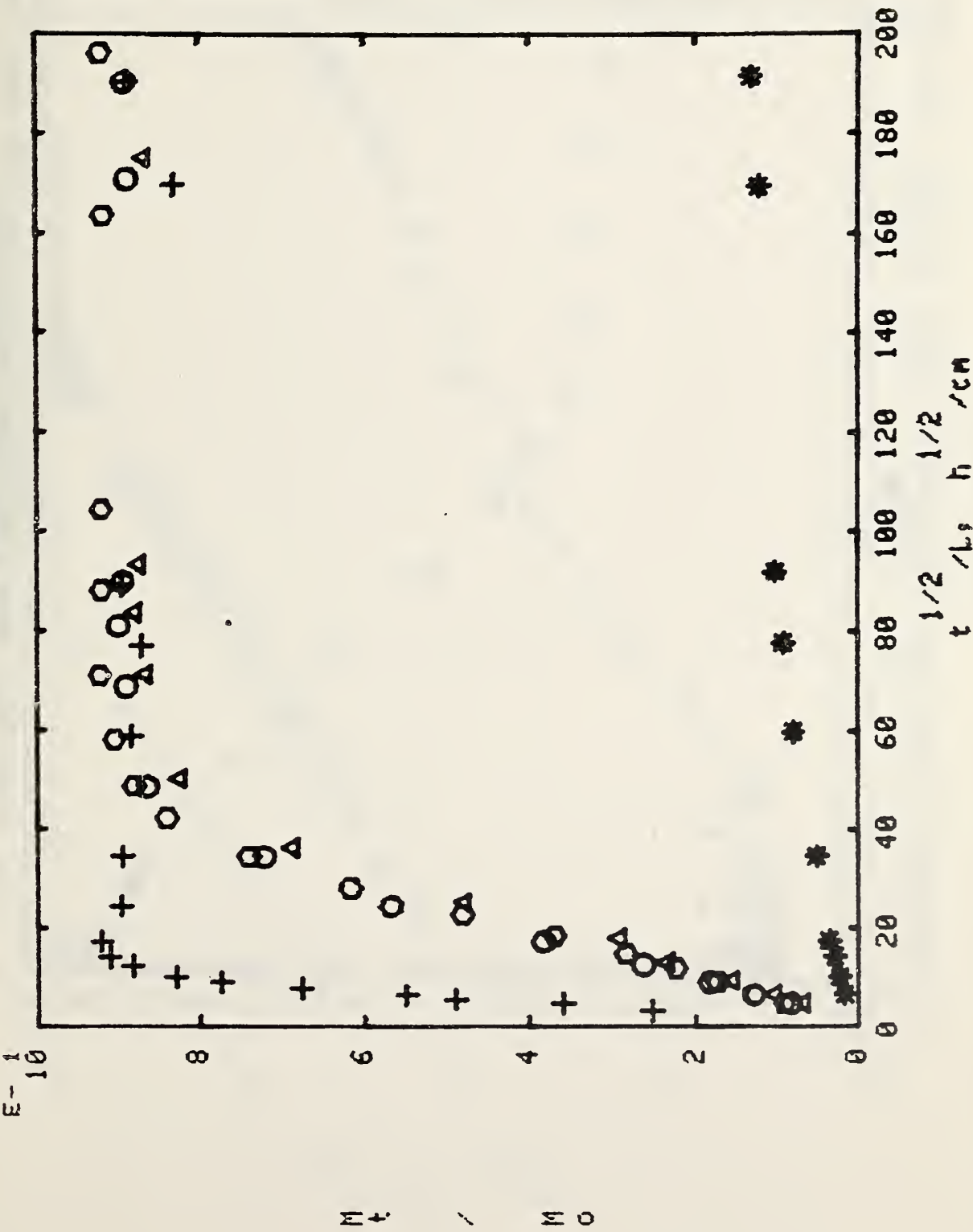


Figure V-1.2-C

MIGRATION OF BHT FROM ETHYLENE-5% VINYL ACETATE COPOLYMER
AT 60°C

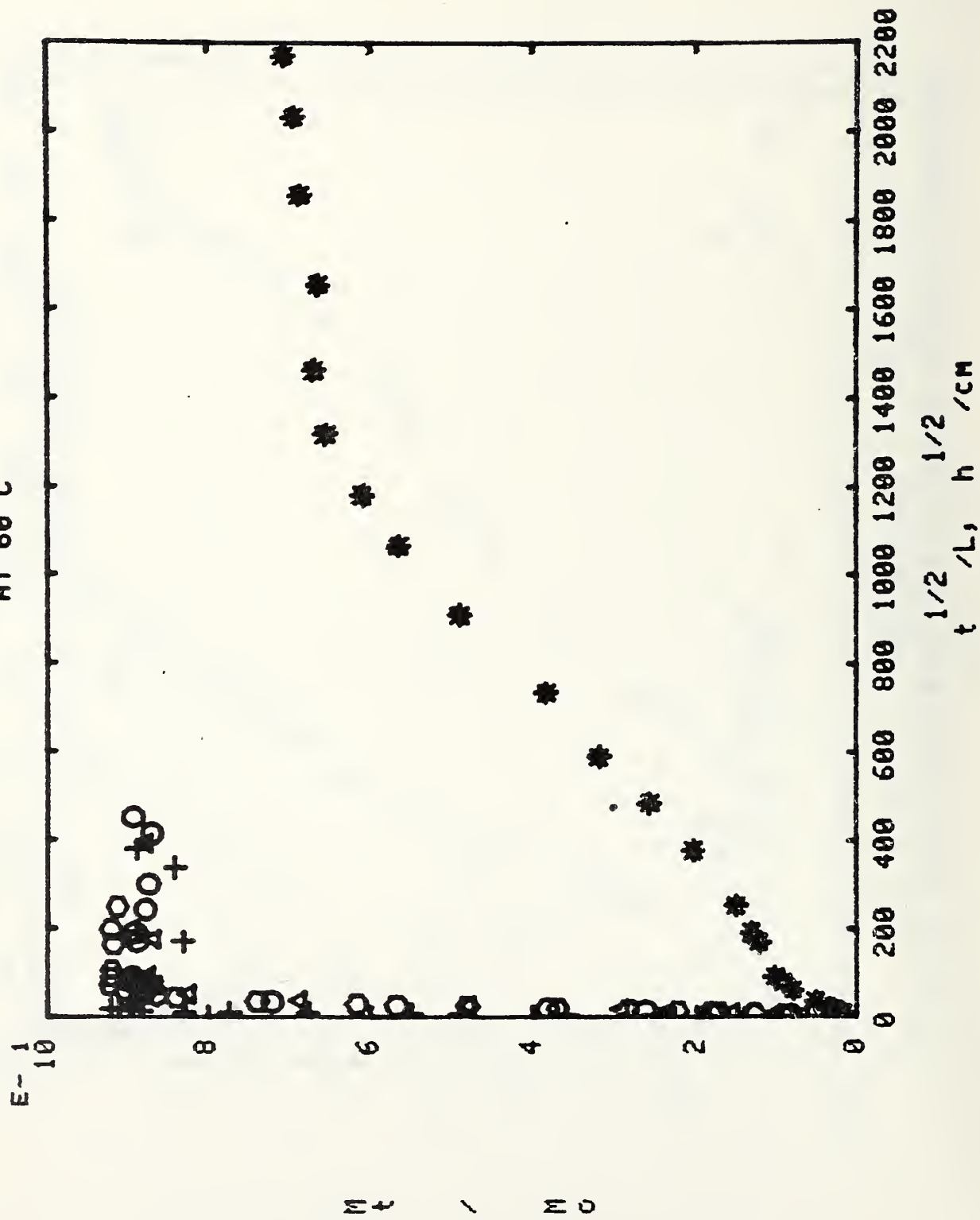


Figure V-1.3-A

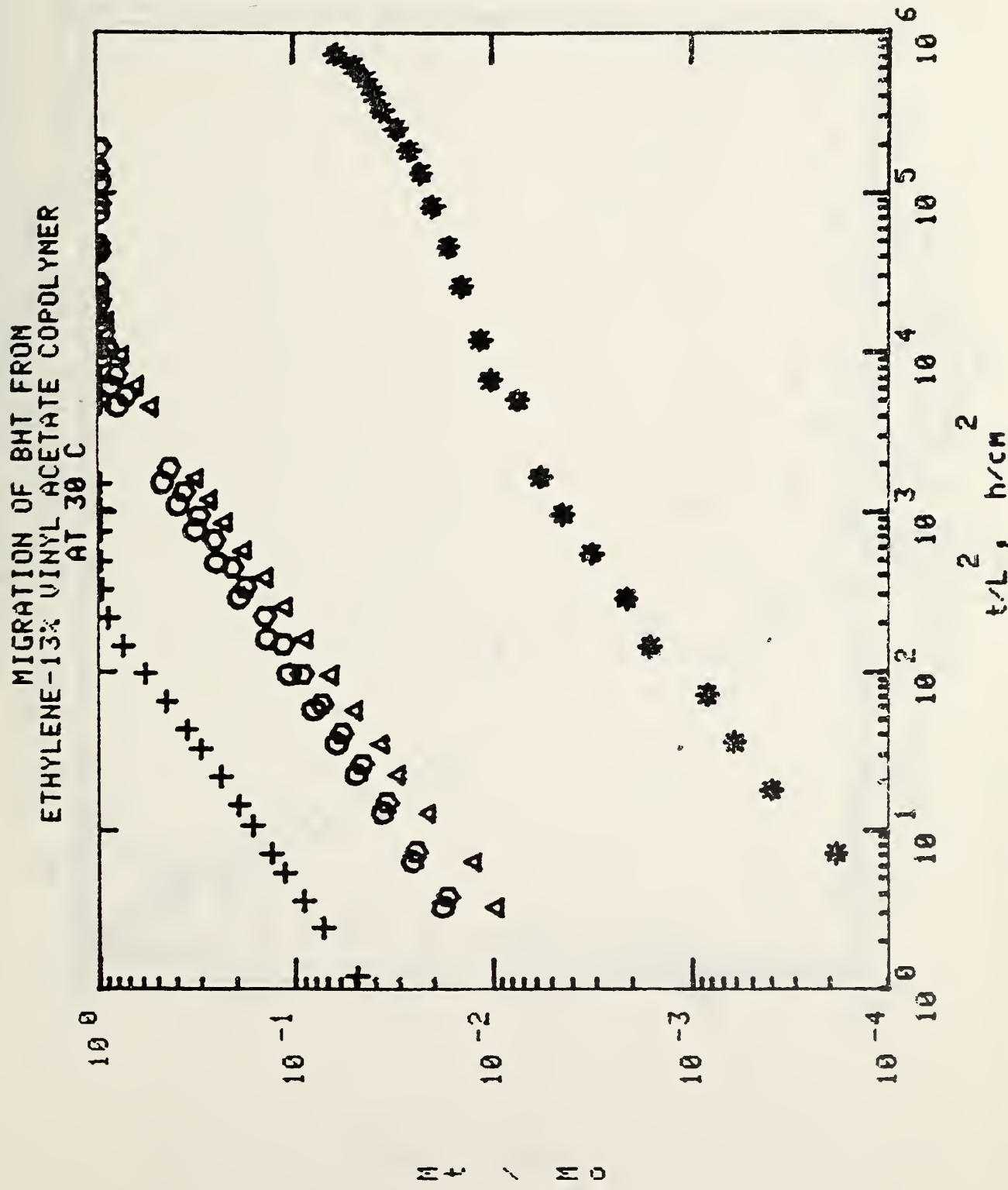


Figure V-1.3-B

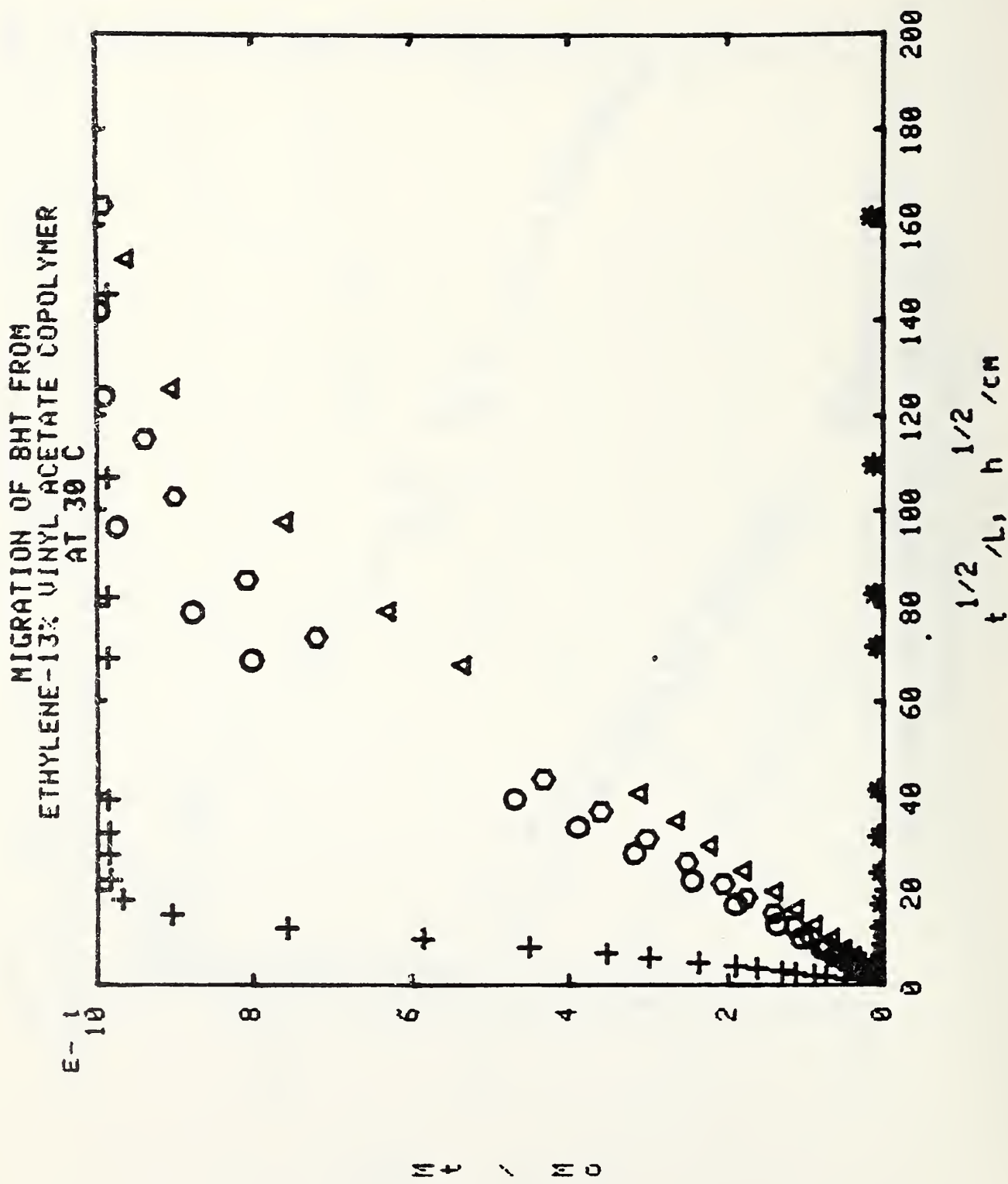


Figure V-1.3-C

Migration of BHT from
Ethylene-15% Vinyl Acetate Copolymer
at 30°C

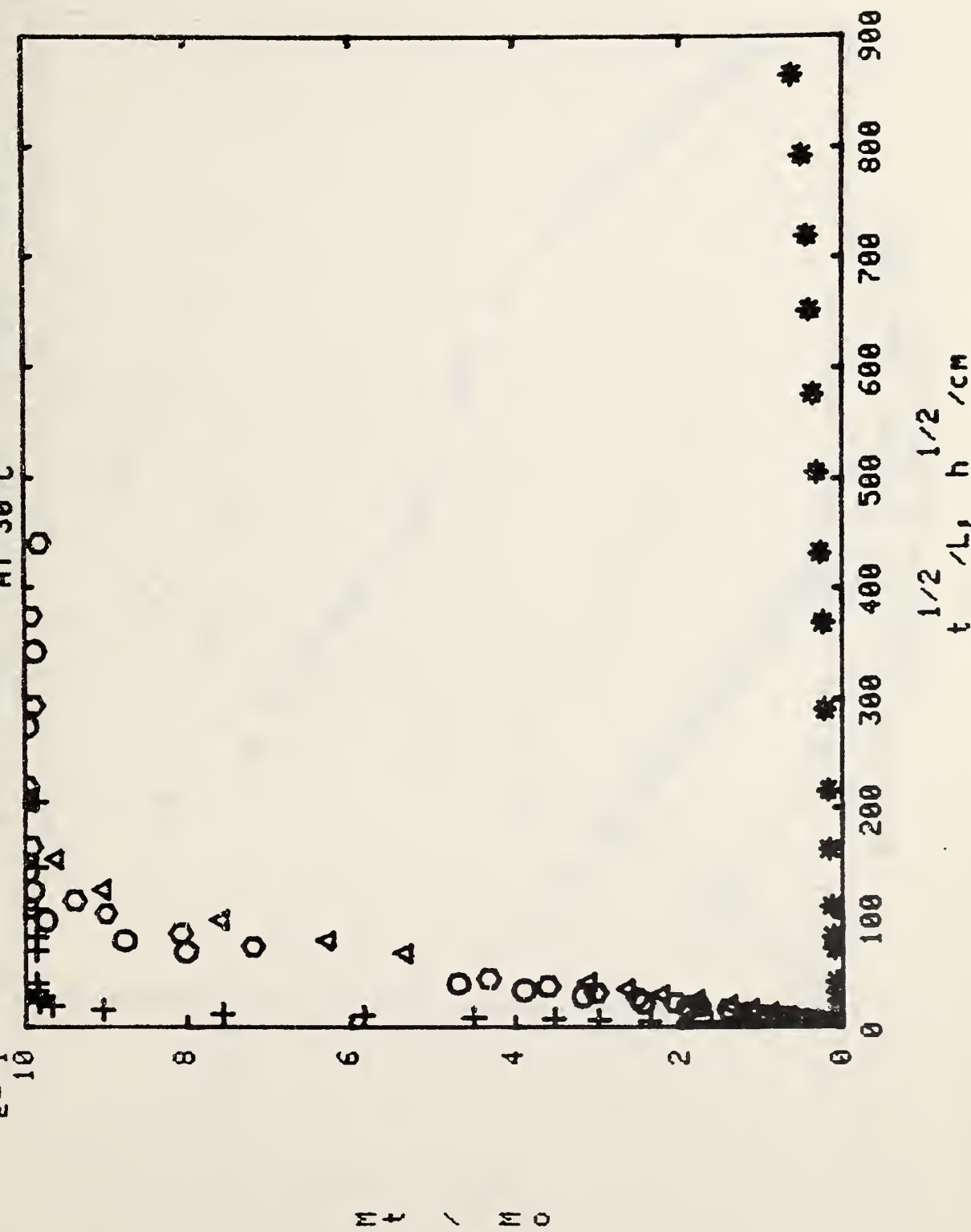


Figure V-1.4-A

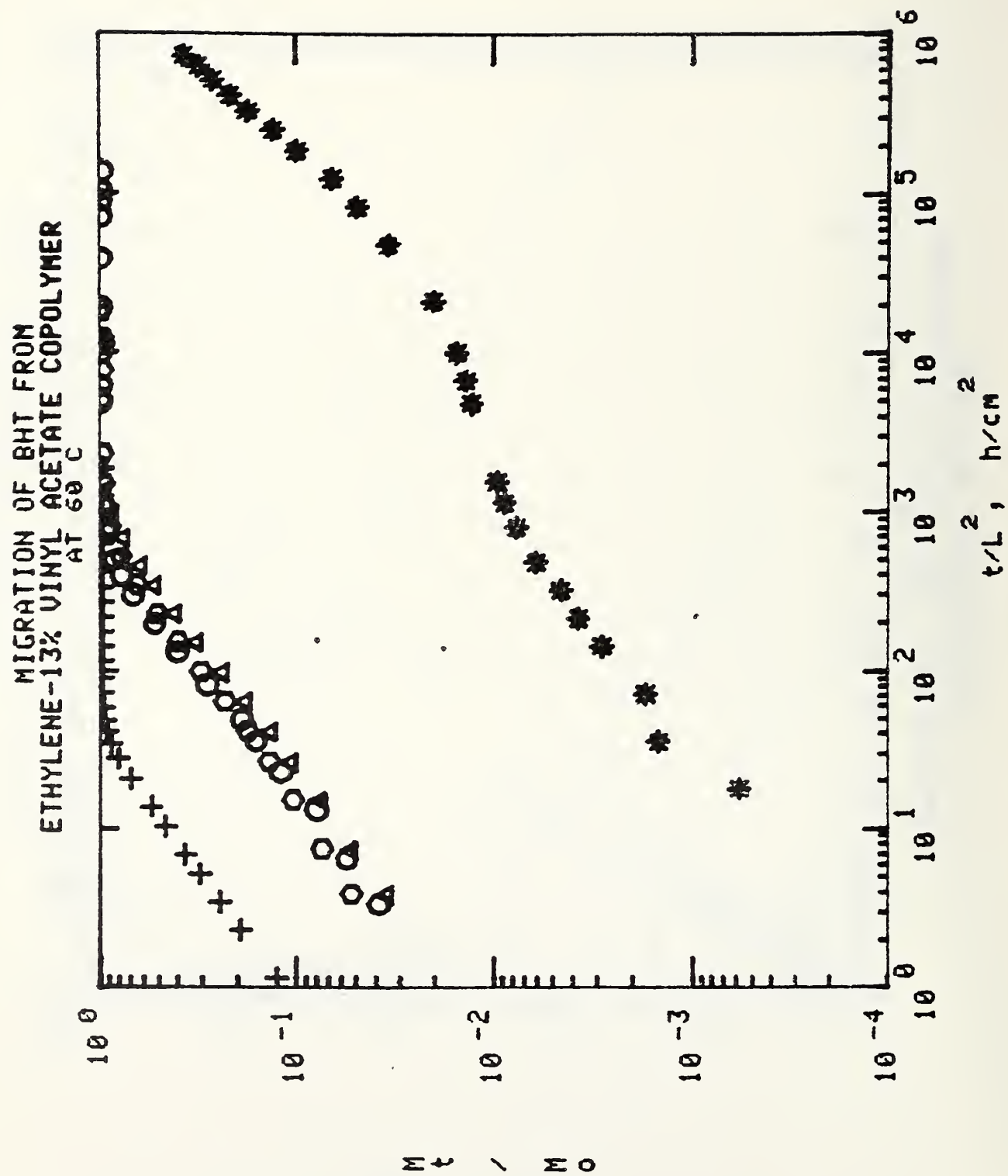


Figure V-1.4-B

MIGRATION OF BHT FROM
ETHYLENE-13% VINYL ACETATE COPOLYMER
AT 60°C

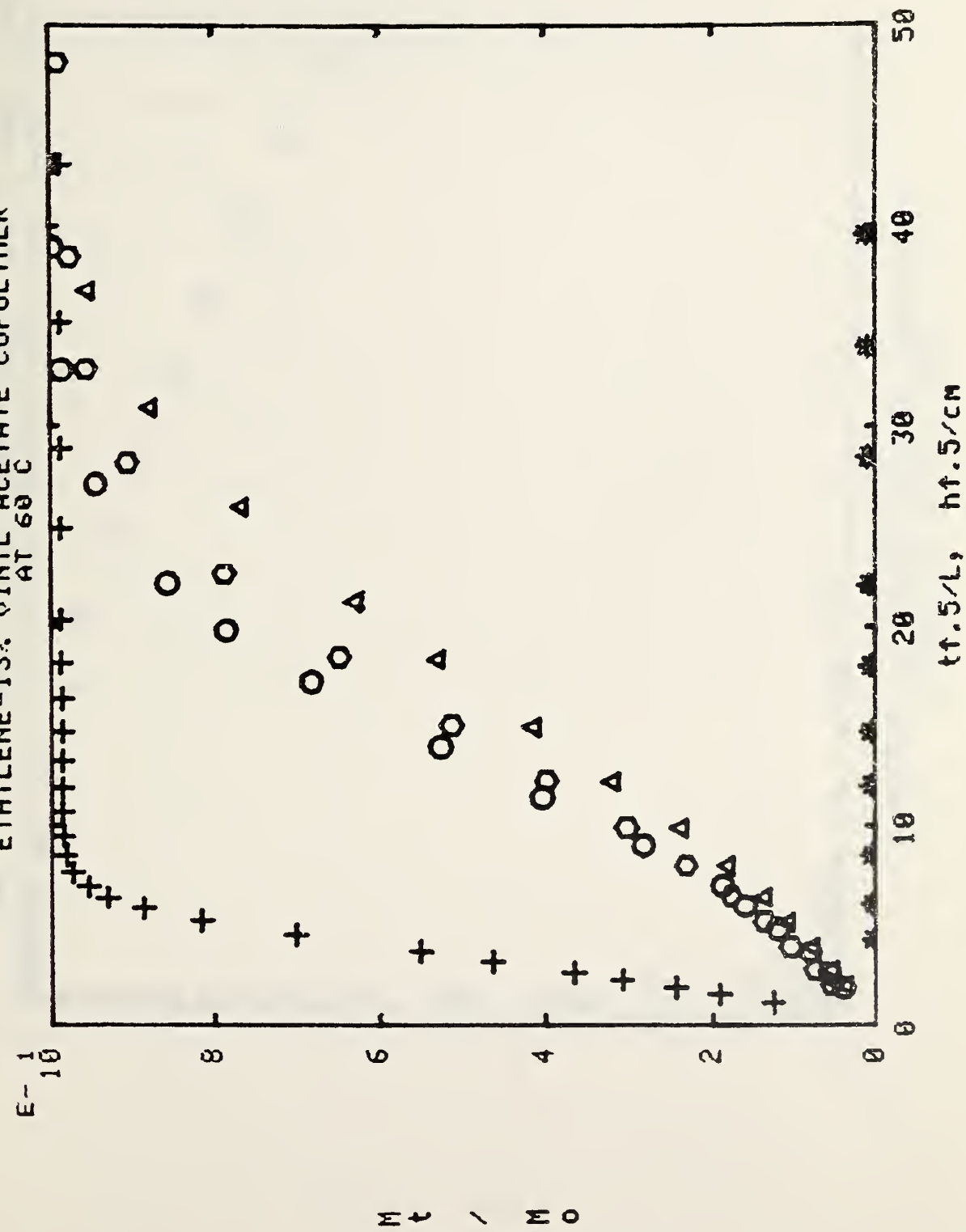


Figure V-1.4-C

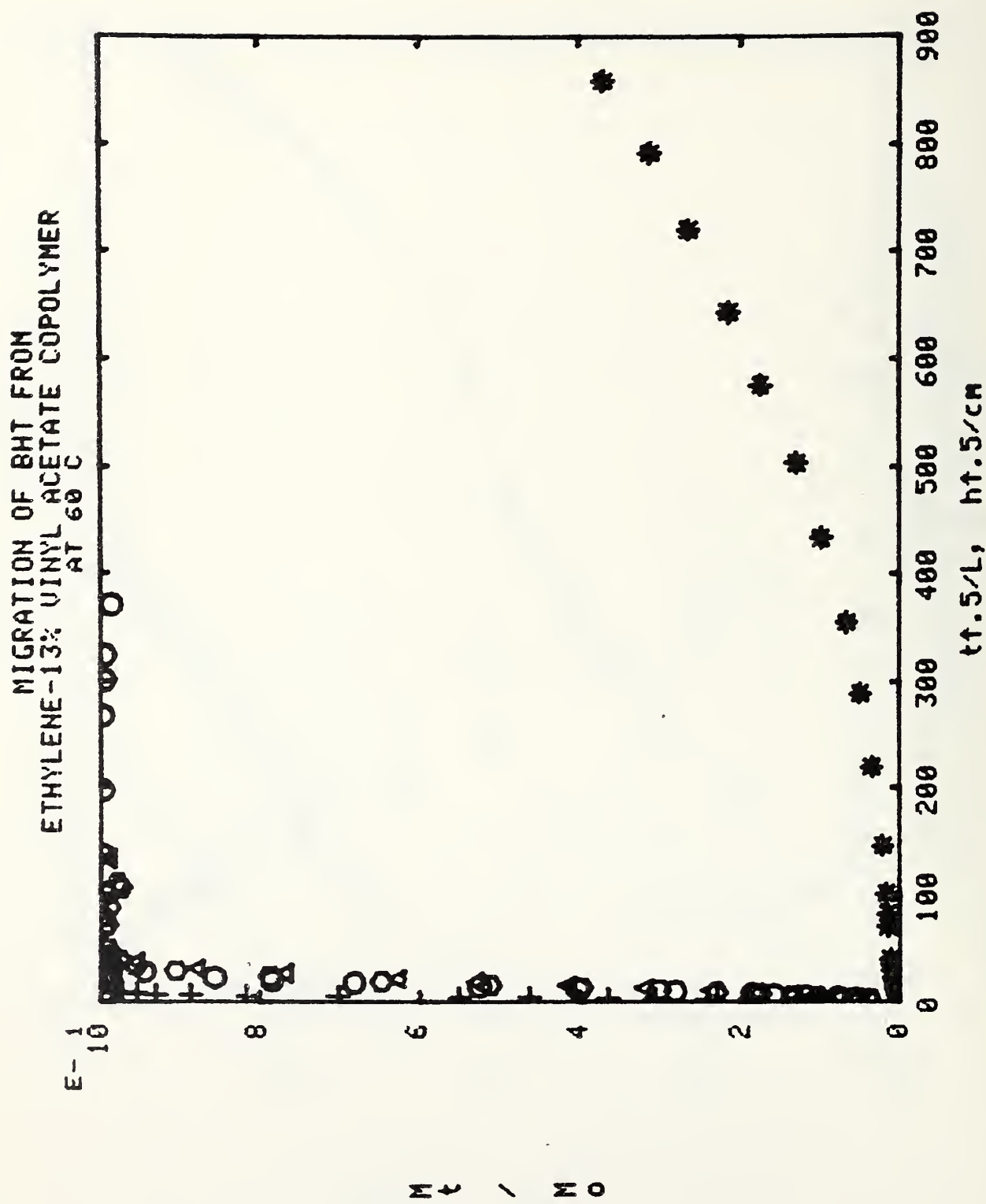


Figure V-2.1 Migration of BHT from E-VA Copolymers into n-Heptane

Figure V-2.2 Migration of BHT from E-VA Copolymers into Corn Oil

Figure V-2.3 Migration of BHT from E-VA Copolymers into Ethanol

Figure V-2.4 Migration of BHT from E-VA Copolymers into 95% Ethanol

Legends for Figure V-2

	E-5% VA	E-13% VA
30°C	+	◇
60°C	x	□

Figure V-2.1

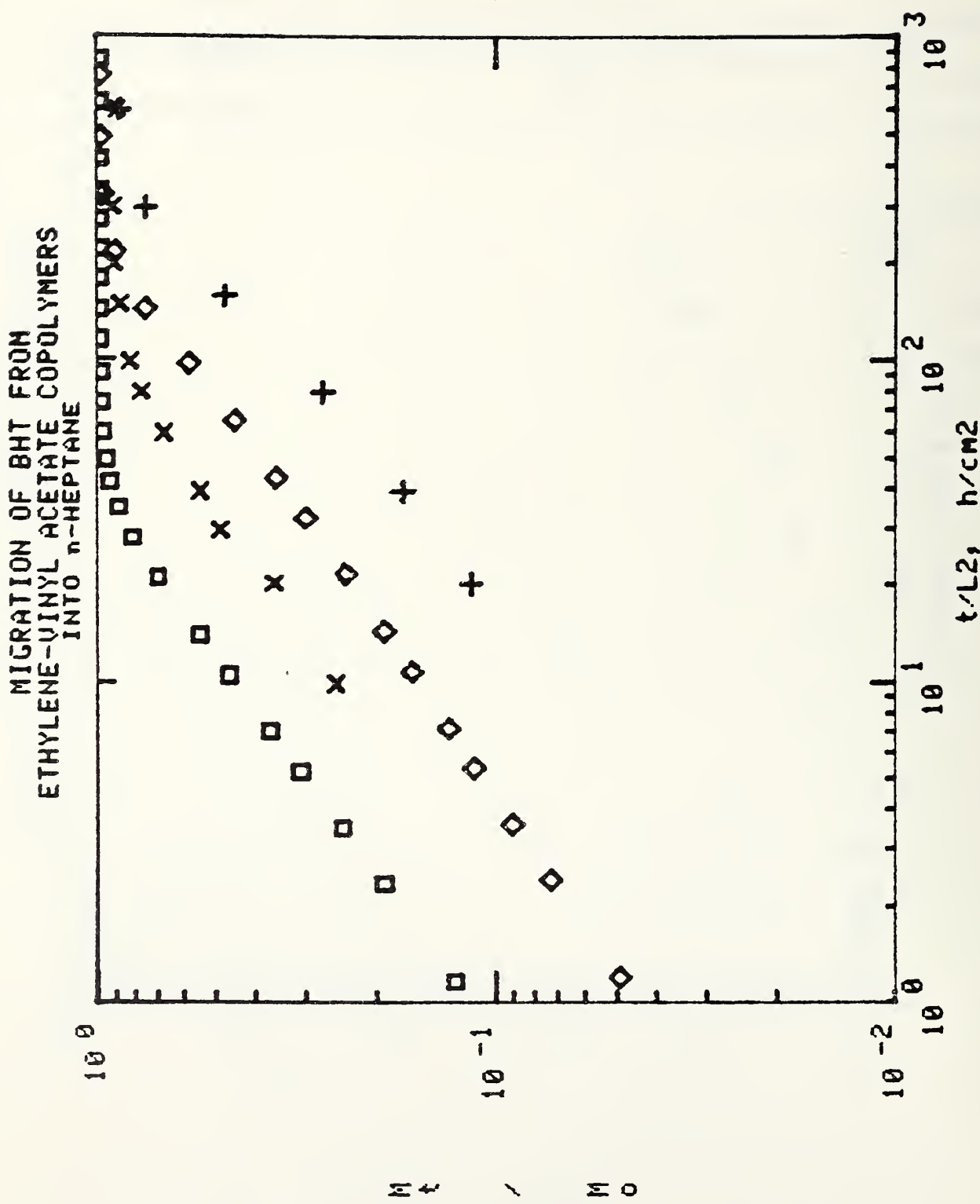


Figure V-2.2

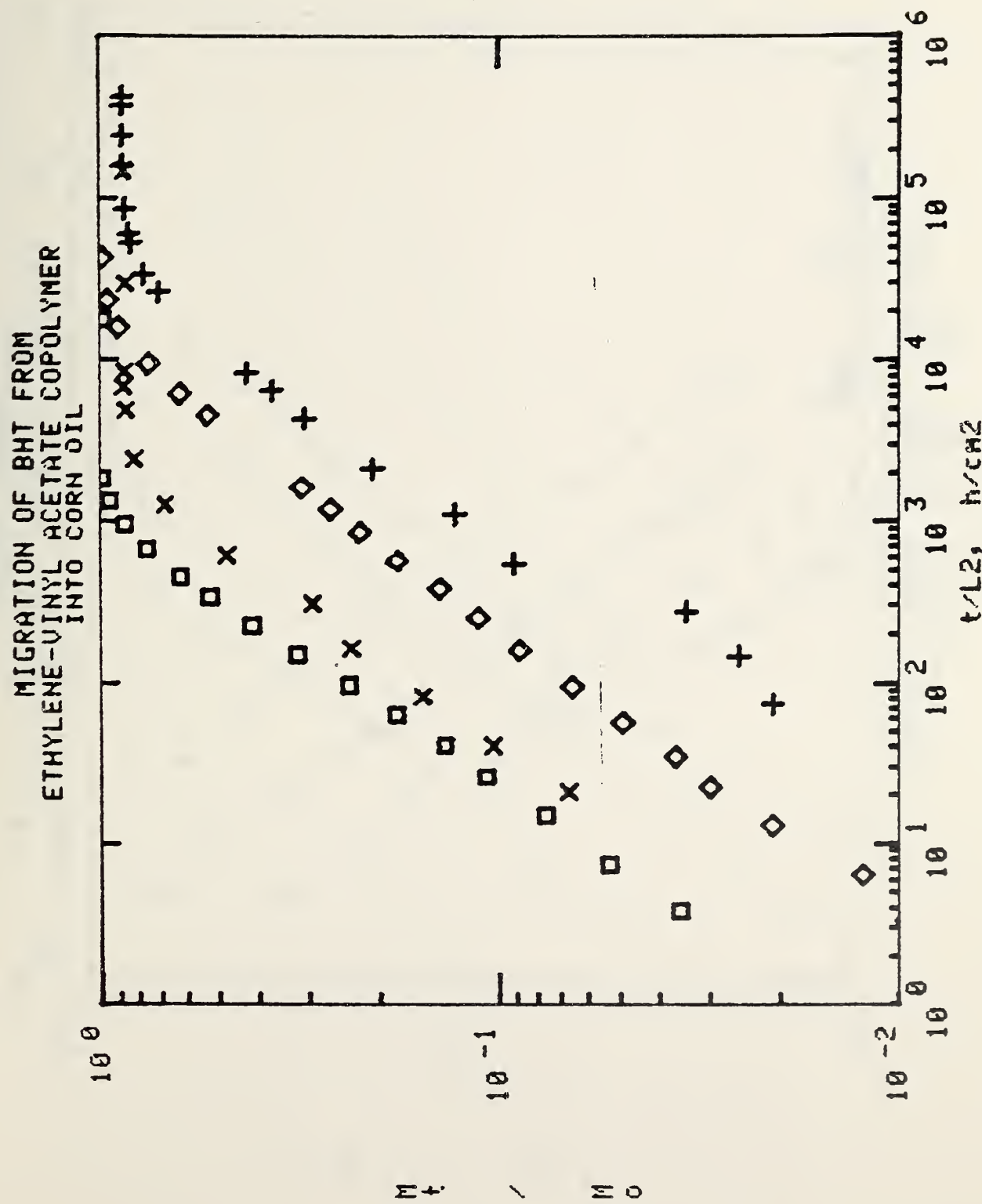


Figure V-2.3

MIGRATION OF BHT FROM
ETHYLENE-VINYL ACETATE COPOLYMER
into ETHANOL

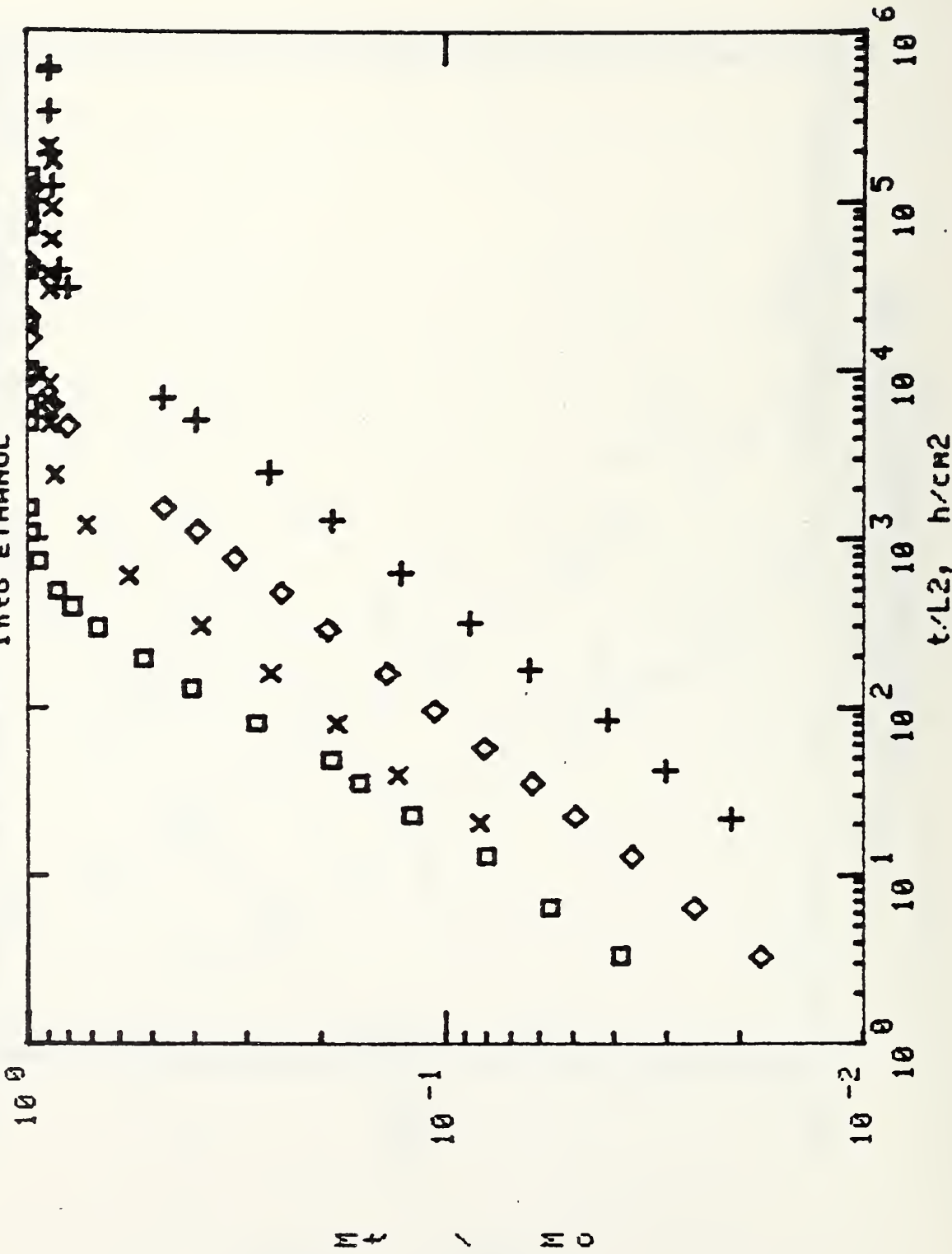


Figure V-2.4

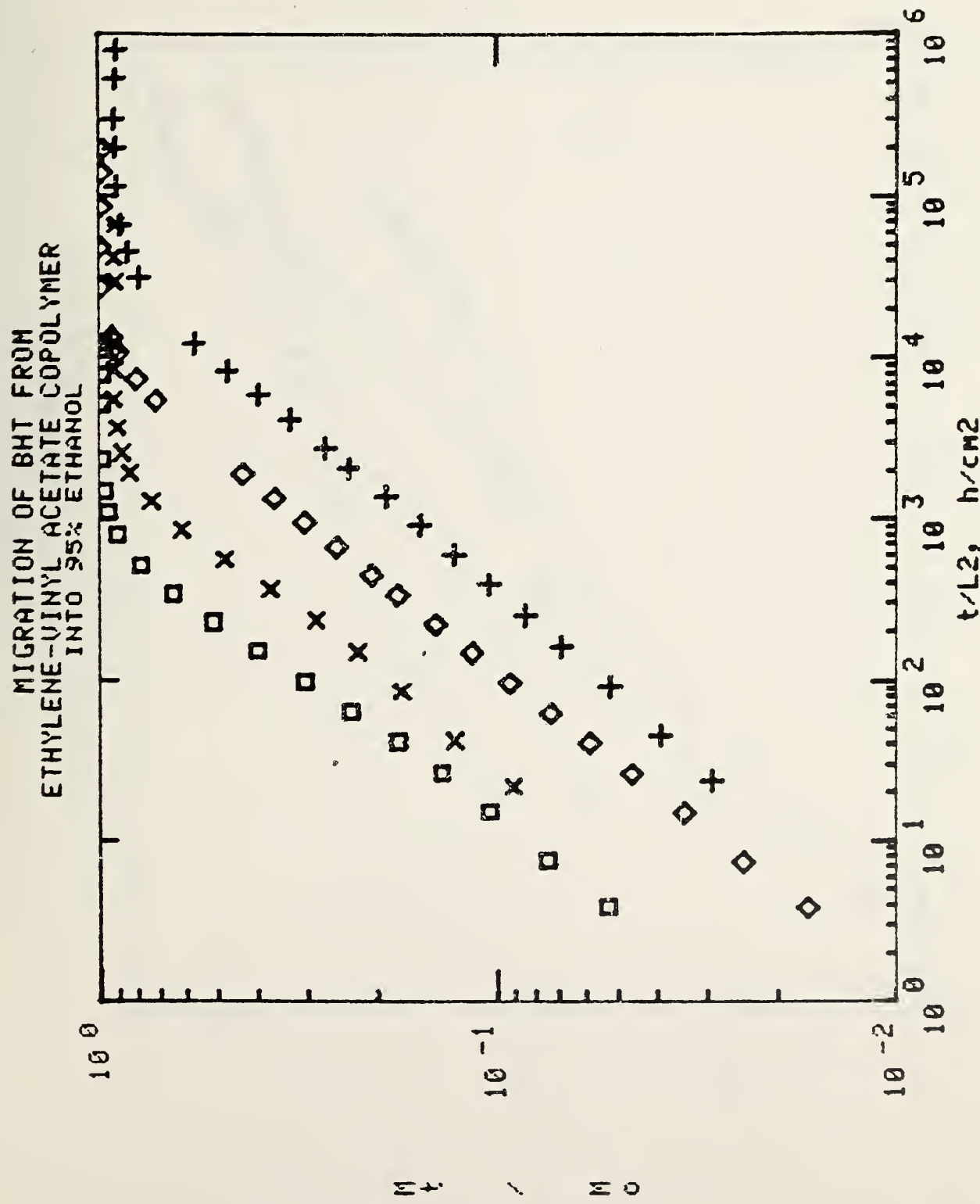


Figure V-3 A, B, C Migration of BHT from E-VA Copolymers into Water

Legends

	E-5% VA	E-13% VA
30°C	+	◇
60°C	x	□

Figure V-3-A

Migration of BHT from
Ethylene-Vinyl Acetate Copolymer
into Water

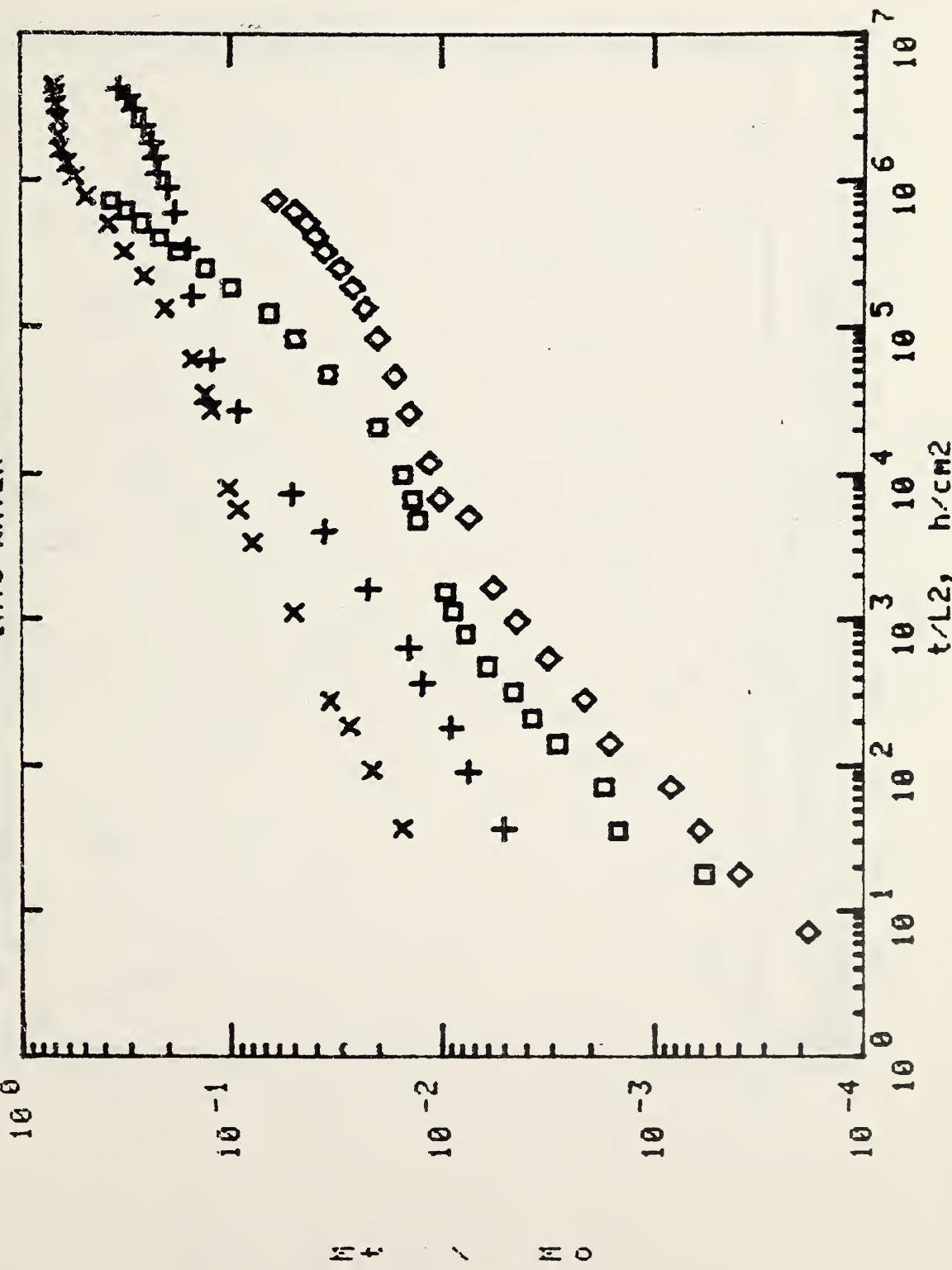


Figure V-3-B

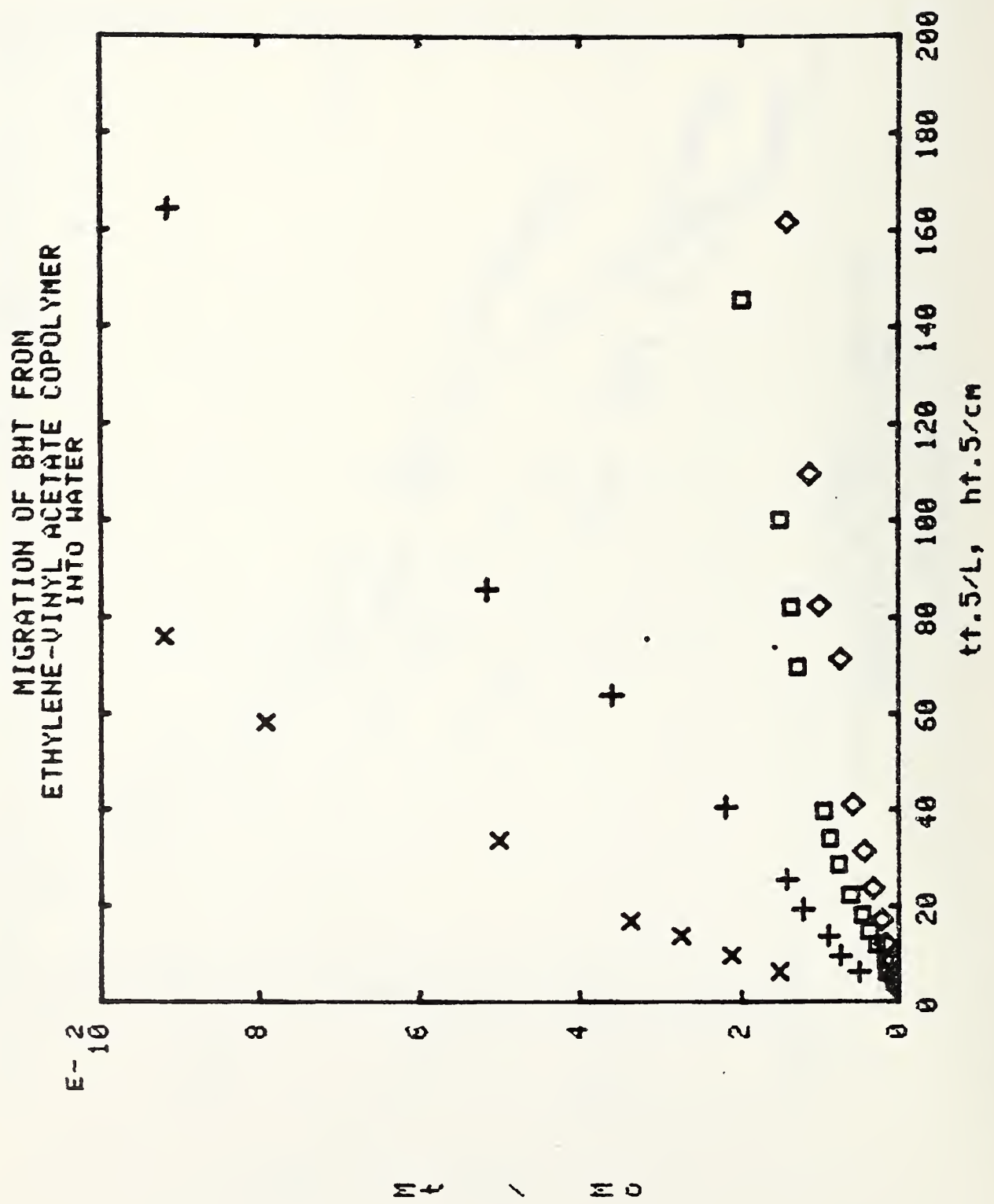
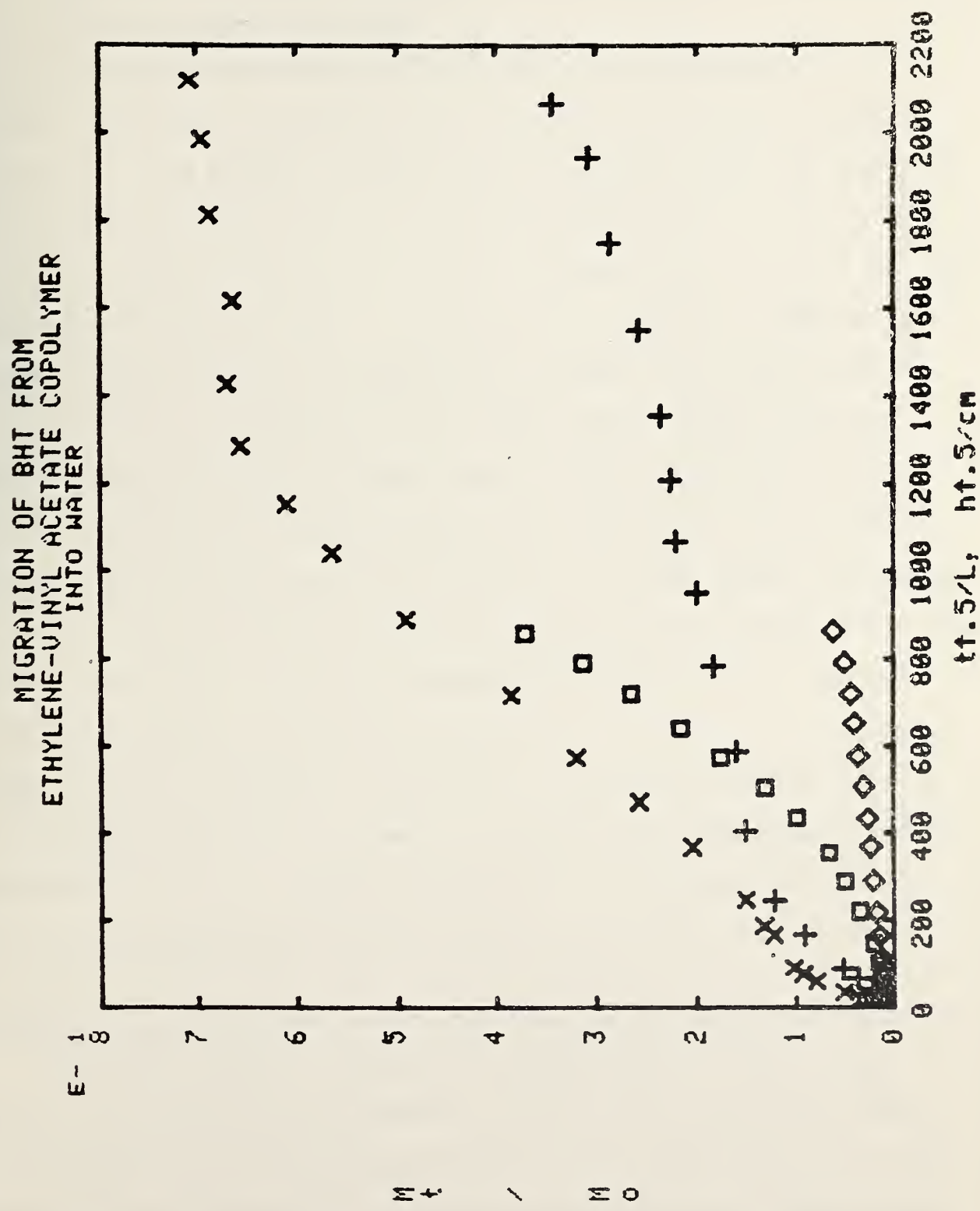
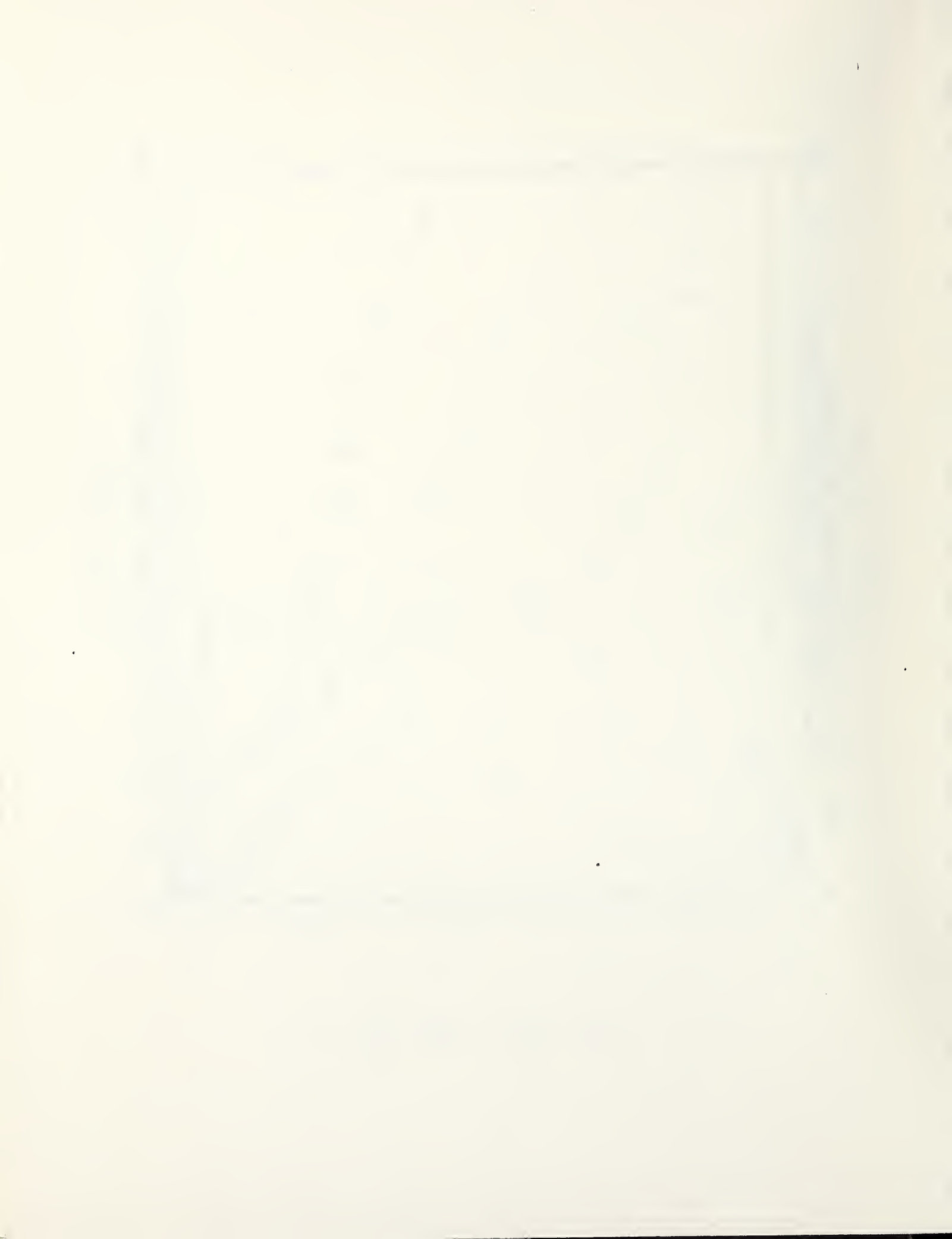


Figure V-3-C





VI. IGC Determinations of the Relative Diffusion Coefficients of Several Probes in Linear Polyethylene

Inverse gas chromatography (IGC) studies on the determination of diffusion constants in polyethylene were reported in the previous annual report [1]. These findings indicate that while accurate absolute diffusivity measurements seem beyond the reach of the IGC method, the possibility exists for using the technique to rank the relative ease of diffusion of many probes in a given polymer. A calibration of a specific column can be accomplished by conducting experiments with a probe-polymer combination of known diffusivity to calculate the mean polymer layer thickness from the measured peak broadening. The same column can then be used under similar operating conditions to determine other probe diffusivities and a relative ranking of many probe diffusion coefficients can be established. Experiments on several series of compounds have been conducted and those for which the data analysis and reduction have been completed will be used to illustrate the procedure of relative diffusivity determinations. Preceding this discussion, however, will be: a review of the theory for determining absolute diffusion constants from IGC studies, a summary of results from polyethylene-octadecane experiments to be used in the relative diffusivity determinations, and a section giving experimental details on the relative diffusivity measurements.

A Review of Pertinent Theory for IGC Absolute Diffusivity Determinations

The probe or migrant injected onto a chromatographic column resides partly in the carrier gas and partially in the polymer or stationary phase during transit through the column. Diffusion of the probe vapor in the gas phase can occur parallel to the background flow of carrier gas sweeping the vapor through the column both in the direction of and opposed to the carrier flow. This longitudinal or axial gaseous diffusion process causes an increase

in width of the eluted peak over its initial injection distribution. An additional gas phase peak spreading mechanism originates in the multiplicity of path lengths possible around the packed support particles and leads to an associated broadened distribution of elution times. Peak spreading can also be attributed to diffusive processes in the stationary phase. If the carrier gas flow rate is very rapid, steady-state partitioning of the vapor between the gas and stationary phases cannot be established. The residence time in the stationary polymer phase is then influenced by both the diffusivity of the probe in the polymer and the distance in the stationary phase over which diffusion must occur, among other factors. The peak broadening processes discussed above are commonly quantified by the convenient construct of the height equivalent to a theoretical plate, H , which is related to the width at half-height, $w_{1/2}$, for an eluted Gaussian-shaped peak as follows:

$$H = \frac{L}{8 \ln(2)} \left(\frac{w_{1/2}}{t_R} \right)^2 \quad (1)$$

where L is the column length, and t_R the retention time of the peak at the maximum. The determination of these quantities from a Gaussian-shaped experimental chromatogram with a peak standard deviation of σ is illustrated in the schematic of Figure VI-1.

An equation attributed to van Deemter et al. [2] is often used to relate H to the local carrier gas flow velocity, u , and is of the form:

$$H = A + B/u + Cu \quad (2)$$

The A term arises from the number of possible gas paths through the column; B is from longitudinal diffusion of the probe in the carrier gas; and C originates from the resistance to mass transfer in the polymer or stationary

phase. The coefficients in equation 2 are defined as follows:

$$A = 4\lambda r \quad (3)$$

$$B = 2\gamma D_g \quad (4)$$

$$C = \frac{8 s^2 k}{\pi^2 D_s (1 + k)^2} \quad (5)$$

where λ is a measure of support packing irregularities, r the average support particle radius, γ a correction coefficient for constriction and tortuosity of the gas flow path in the column, D_g the probe diffusivity in the gas phase, s the effective thickness of the stationary phase film, D_s the diffusivity of the probe in the stationary phase, and k the column capacity factor commonly given as:

$$k = \frac{t_R - t_M}{t_M} \quad (6)$$

where t_M is the time taken to elute an unsorbed marker material from the column. Equation 2 indicates that, at large values of the flow rate, H will be proportional to the flow rate. The probe-polymer diffusion coefficient can then be determined from the constant of proportionality C by equation 5 if the effective film thickness is known. Equation 5 is not applicable for columns packed with glass bead support particles. In such cases of diffusion into a uniform stationary phase film on a solid surface, a similar relation for C applies:

$$C = \frac{2 s^2 k}{3 D_s (1 + k)^2} \quad (7)$$

While equation 2 gives a suitable explanation of peak spreading in some cases, other results have raised questions about its utility, particularly when significant gas phase diffusion can occur. These discrepancies indicate that the simplifications made by van Deemter in neglecting other mechanisms of diffusion are not valid for all systems. Giddings and co-workers [3] examined several modified forms of the van Deemter equation and found that none could account for the experimental results obtained on columns of varying length and outlet pressure.

An equation proposed by Jones [4] takes several additional sources of peak spreading into consideration: resistance to mass transfer in the gas phase normal to the carrier gas flow direction (D), carrier gas velocity distribution effects (E), and a correlation between these two processes (F). This equation is given as:

$$H = A + B/u + (C + D + E + F)u \quad (8)$$

where C is as in equation 7 and

$$D = \frac{C_D k^2 g^2}{D_g (1 + k)^2} \quad (9)$$

$$E = \frac{4C_E r^2}{D_g} \quad (10)$$

$$F = \frac{4\theta(C_D C_E)^{1/2} k r g}{D_g (1 + k)} \quad (11)$$

g being the diffusion path length in the gas, θ the correlation coefficient between the D and E terms, and C_D and C_E geometric constants.

The radial diffusion D term was discounted by van Deemter and co-workers in the derivation of equation 2 because of the much greater magnitude of D_g compared to D_s . This simplification is valid for the thick liquid films used in their work, studies of columns with roughly 0.3 mass fraction stationary phase loading. The diffusion distances s and g are roughly comparable under these circumstances. If low column loadings which reduce s to about 0.01g are used, then the much larger probe diffusivity in the gas phase is overcome by the ease of mass transfer through the thinly-spread liquid phase and the D term cannot be ignored.

The E contribution to spreading arises from diffusion caused by differences in the velocity of the carrier gas in various parts of the column. Velocity differences can arise from both short and longer range channels formed by support packing irregularities and from trans-column effects caused by the influence of the tube walls. These effects are summarized by Giddings [5] who estimates approximate magnitudes for contributions of the many possible velocity gradients combined in the E term of equation 8.

While the first three terms in equation 8 are independent of one another, the D and E terms are not since molecules which have a lower average velocity will spend a smaller fraction of time in the gas phase and therefore not experience as great a percentage of gaseous diffusion as molecules with higher than average velocity. The total variance, σ_{ij}^2 , of two such dependent contributors, σ_i^2 and σ_j^2 , is given as:

$$\sigma_{ij}^2 = \sigma_i^2 + \sigma_j^2 + 2\theta_{ij}\sigma_i\sigma_j \quad (12)$$

where the correlation coefficient between the two contributors is denoted as θ . The F term of equation 11 originates from the final term in the above relation and corresponds to the interaction between the D and E terms.

Giddings [6] has given an expression said to account for all known plate height terms contributing at least one percent of the total plate height for packed GC columns in the absence of interfacial adsorption as:

$$H = \sum_i \frac{1}{1/A_i + 1/E_i u} + \frac{B}{u} + \sum_i C_i u + Du + G \quad (13)$$

The G term represents the contribution of column-wide velocity inequality effects, such as that arising from column coiling, and could alternatively be included in E. This equation is otherwise similar to equation 8 except for its omission of the interaction term F. A compilation of the C, D, and E coefficients expected to apply in various situations and a means of calculating their magnitude from measurable column parameters is given in reference [5].

The plate height equations discussed up to this point apply only to the peak spreading in a short region of the chromatographic column where the carrier velocity u is a constant local to that region. A final modification can be made to equation 8 or 13 to account for the variation in local carrier velocity along the column caused by the drop in pressure from column inlet to outlet. The effect of this correction is primarily on the liquid mass transfer term, as discussed by Dal Nogare and Juvet [7]. The final expression for the dependence of the experimentally accessible column average plate height, \bar{H} , on the carrier gas outlet velocity, u_o , is similar to equation 8 and is given as:

$$\bar{H} = A + \frac{B}{u_o} + \left(\frac{2p_o C}{p_i + p_o} + D + E + F + G \right) u_o \quad (14)$$

where p_i and p_o are the column inlet and outlet pressure. The gas phase diffusivity found in equations 4, 9, 10, and 11 must also be restated at the outlet pressure of the column. The form of this modification is based on the relationship between the local column velocity and the average carrier gas velocity. The preceding treatment assumes a distance average of this velocity. Other methods [3] involve the use of a pressure average velocity and yield a slightly different equation for \bar{H} but agree reasonably well with equation 14 even at large values of the inlet pressure.

The magnitudes of coefficients in equation 14 and the configuration of the polymer film on the surface of the supporting beads can have a significant influence on the methods used in interpreting data derived from IGC determinations of diffusive peak broadening as discussed in the previous annual report. These effects will be reviewed briefly for the experimental determination of the diffusivity of normal octadecane in high density polyethylene considered there.

Summary of Previous Results for LPE-Octadecane

As discussed earlier, IGC determinations of diffusivity in the polymer phase are made from a study of probe peak broadening at varying carrier gas velocities; a schematic illustration of the results is shown in Figure VI-2. The local plate height and carrier velocity of equation 2 are commonly replaced by the measured average quantities in experimental determinations of column efficiency. Such \bar{H} vs \bar{u} plots for eight flow velocities in the flow rate region from about 10 to 100 cm/s yielded a slope C for column A of 7.05 ms with calculated standard deviation of 8.8 percent and 18.9 ms with calculated standard deviation of 7.4 percent for B. The van Deemter treatment provides for the calculation of the diffusivity of C_{18} in linear

Polyethylene from these C values by equation 7 when the polymer film thickness in each column is known. Such a procedure is proper only if the other contributions to peak spreading discussed earlier are small when compared to that arising from polymer or stationary phase mass transfer. This assumption has been made in many previous studies of probe-polymer diffusion. However, if the other gas phase kinetic processes are operating to a significant extent then the limiting slope at high flow rate of Figure VI-1 should also reflect these contributions, as indicated by equations 8 and 14. It was important, therefore, to estimate the relative magnitude of the gas phase terms as compared to the stationary phase contribution to determine if such a simplified data analysis procedure could be applicable.

A further consideration is the nature of the carrier velocity on which the plate height depends. Equation 14 indicates that while the plate height contribution of the B, D, E, and F terms are directly or inversely proportional to the carrier outlet velocity u_o , the C term contribution to \bar{H} depends on the average velocity. Consequently, the often-employed plot of \bar{H} vs average carrier velocity suggested by equation 2 will not be linear at high flow velocity for any variant of velocity if both stationary and gas phase mass transfer effects are nonnegligible.

The estimated gas phase contributions to peak spreading were computed for both columns and compared to the limiting slope C calculated from the van Deemter equation least squares fit. It was evident that the peak spreading attributed to gas phase diffusion made a significant contribution to the overall kinetic broadening for column A. In instances of this nature, where gas phase diffusion terms make a significant contribution to peak spreading, difficulty can be encountered in deciding which are important and calculating their values. Alternatively, it is possible to resolve the gas and stationary

phase mass transfer contributions without directly estimating the magnitude of the gas phase terms. Purnell [8] has suggested a method which involves first determining A and B of equation 14 and employing these values to reduce the experimental \bar{H} to the peak spreading arising only from mass transfer effects, \bar{H}' . In terms of an equation:

$$\bar{H}'/u_0 = \left(\bar{H} - A - \frac{B}{u_0} \right) / u_0 = \frac{2p_0}{p_i + p_0} C + D + E + F + G \quad (15)$$

When the left-hand side of this equation is plotted against the pressure correction $2p_0/(p_i + p_0)$, a straight line with slope C and intercept $(D + E + F + G)$ should be obtained. Evaluation of the constants A and B requires that experiments be conducted over a wide range of carrier gas flow velocities. Limited low velocity data often precludes experimental determination of these constants in many cases; however, they can be estimated from theoretical considerations and experimental results. The multipath A term defined in equation 3 has been determined to have a λ value between one half and unity in many cases [9] and a value of 0.5 was assumed here. The longitudinal diffusion B term most often has a γ value near one [9] and unity was utilized in the determinations. These values were used to calculate the A and B terms of equation 19 and a fit of \bar{H}'/u_0 against the pressure correction term gave a linear least squares line with a slope of 16.4 ms and calculated standard deviation of 5.5 percent for column A and 21.9 ms with calculated standard deviation of 6.6 percent for B. These results are compared to the van Deemter fit slopes in Table VI-1 on the following page. The smaller standard deviations of the Purnell method slopes compared to the corresponding values for the van Deemter fit discussed earlier indicated the better applicability of the Purnell method. The C value obtained by this method for column A was more

Table VI-1

Comparison of the C Coefficient Determined by Two Methods
for LPE-Octadecane at 150 °C

Column Code	Column Loading (Mass Fraction Polymer)	$C_{\text{van Deemter}}$ (Eq. 2)	C_{Purnell} (Eq. 15)
A	6.10×10^{-4}	7.05 ms	16.4 ms
B	5.71×10^{-3}	18.9 ms	21.9 ms

than twice that obtained from the van Deemter analysis. This was attributed to the latter's combination of the effects of polymer and gas phase mass transfer in a nonlinear manner, which can be noted by comparing the C term appearing in equation 2 to its complement in equation 14, the expression in parenthesis. For the case of column A where the gas phase terms contributed significantly to the peak broadening, the van Deemter equation-based analysis underestimated the magnitude of C. Gas phase mass transfer effects for column B contributed only slightly to the total broadening and this was reflected in the similar results of the two methods for determining C which differed by only 15 percent, as indicated in Table VI-1.

While the method of determining the polymer phase mass transfer contribution can be important under certain column conditions, the most significant parameter which affected the diffusivity determination was the distance over which diffusion occurred. This distance is commonly identified with the effective film thickness s of equation 7. It is often assumed that the stationary phase is distributed as a uniform thin film over the entire surface of the spherical glass bead column packing. The effective film thickness for this geometry can be calculated from the expression:

$$s = \frac{M_s}{4\pi r^2 N p_s} \quad (16)$$

where M_s is the mass of polymer phase in the column, N the number of support particles in the column, and p_s the polymer density. Such a configuration requires a high amount of stationary phase surface area, however, and a geometry in which the stationary phase is confined by capillary forces into annular regions around the contact points between support particle spheres has been advocated as a means of minimizing the exposed stationary phase surface area. This geometry has been observed for both low molecular weight [6] and polymeric [10] stationary phases coated onto glass bead supports and is illustrated schematically in Figure VI-3. A stationary phase film may also be present on surfaces where the beads are not in contact, as indicated in the inset. Its thickness, however, should be much less than that of the stationary phase in the region of adjacent support beads.

The different diffusion path lengths expected for the two extremely different polymer distributions described above can exert a significant influence on determinations of polymer-probe diffusivity by IGC. Giddings [11] has compared the contribution to peak spreading expected from a hypothetical support coated with two regions, a thin film of uniformly adsorbed stationary phase and a second region where the liquid stationary phase collects at the support contact points. He concludes that for glass bead supports at mass fraction loadings up to about 0.015, the bulk of the stationary phase is present at the bead contact points and controls the plate height performance by virtue of its greater thickness. An equation has been proposed [12] to relate C to measurable column parameters for this geometry and is as follows:

$$C = \frac{r^2 k}{3 D_s (1 + k)^2} \left(\frac{m_s \rho}{18.75 \rho_s} \right)^{1/2} \quad (17)$$

where ρ is the density of the support particles and m_s the mass fraction of polymer phase in the column. This treatment assumes that the spherical support particles pack randomly and are surrounded by 6.25 nearest neighbors. The relation for C is similar in form to equation 7. If the differing variables are combined into an explicit expression for s in terms of the particle size, the following relation is obtained:

$$s = \left(\frac{m_s \rho}{18.75 \rho_s} \right)^{1/4} \frac{r}{2\sqrt{5}} \quad (18)$$

This equation provides for the calculation of a value of s which, when substituted into equation 7, yields a result for C identical to the C term determined by equation 17.

A computation of s by these methods was useful for comparing the effective film thicknesses of the two stationary phase arrangements considered. The average film thicknesses calculated in the above manner for a polymer stationary phase collected at bead contact points were much greater than the thicknesses corresponding to the uniform layer geometry, as can be noted from the following table.

Table VI-2

Effective Layer Thicknesses for Columns with
Differing Polymer Phase Geometry

Column Code	Effective Layer Thickness, $s(\mu\text{m})$ Uniform Thin Surface Film (Eq. 16)	Annular Film at Contact Points (Eq. 18)
A	0.043	5.58
B	0.404	9.77

Both equations 7 and 17 were used to calculate the diffusion coefficient of octadecane in HDPE. For column A assuming a thin uniform film, D_s was calculated from the C coefficient obtained with the Purnell treatment by substituting equation 16 into equation 7; a result of $3.8 \times 10^{-11} \text{ cm}^2/\text{s}$ was obtained. Assuming a geometry where the polymer collects at the bead contact points for column A, D_s was calculated according to equation 17 and a value of $6.4 \times 10^{-7} \text{ cm}^2/\text{s}$ determined. These results and the corresponding values for column B are summarized in Table VI-3 below.

Table VI-3

Influence of Film Geometry on the Diffusion Coefficient
of Octadecane in Linear Polyethylene at 150 °C

Column Code	Diffusion Coefficient ($\text{cm}^2 \cdot \text{s}^{-1}$) Uniform Thin Surface Film (Eqs. 7 and 16)	Annular Film at Contact Points (Eq. 17)
A	3.8×10^{-11}	6.4×10^{-7}
B	4.1×10^{-10}	2.4×10^{-7}

It was evident that the uniform thin film geometry assumption led to unreasonably low values of the diffusion coefficient. In addition, an order of magnitude discrepancy existed between results for the two columns. The assumption of a collection of the polymer stationary phase at bead contact points yielded diffusion coefficients of reasonable magnitude for the temperature of the experiment and column-to-column agreement within a factor of three. This threefold difference in the diffusivity results may be attributed to the effects of differing probe concentrations in the polymer phase of the two columns.

Diffusivity values are often quite sensitive to the concentration of the diffusing species, depending approximately either linearly or exponentially

on concentration [13]. In these IGC determinations the probe concentration in the polymer film varied both along the column length due to increasing peak broadening with transit time and with the depth of penetration into the polymer film expected from Fickian diffusion. The diffusivity determined was therefore a concentration-averaged value over all column conditions. The actual concentration of octadecane in the polyethylene film was estimated to range from 0.01 to 1.5 percent by mass for column A and from 0.001 to 0.16 percent by mass for column B. The diffusion coefficient calculated from the column B results corresponded to probe concentrations about an order of magnitude lower than for column A. This could account for the lower value of probe diffusivity determined with column B. For either a linear or exponential dependence of the diffusion coefficient on concentration, the diffusivity should be less at lower diffusant concentrations, as found here. A determination of the functional relationship of the diffusivity on diffusant concentration was not possible, however, because of the variation in probe concentration along the column length and the limited amount of data available.

Limited data are available in the literature on diffusion of the higher alkanes in LPE; however, studies of similar diffusants, linear aliphatic esters with 25 to 45 backbone units, have been conducted by Klein and Briscoe [14]. Extrapolating their experimental results to an 18 unit chain leads to an expected diffusion coefficient of $5 \times 10^{-6} \text{ cm}^2/\text{s}$ in linear polyethylene at 152 °C. This result is about an order of magnitude greater than the IGC determinations for a polymer film collected at the support bead contact points.

The distribution of film thickness also plays an important role in determining the magnitude of the diffusivity. The previous calculations

assumed that the film thickness was constant. In an actual column, however, the film thickness would be expected to be distributed about the mean values employed here. An alternate form of equation 7 can be used to estimate the effects of different film thicknesses [15]. The equation is:

$$D_s = \frac{k}{C(1+k)^2} \sum_i q_i x_i s_i^2 \quad (19)$$

where q_i is a configuration factor and x_i the volume fraction per unit column volume of stationary phase segments with thickness s_i . Assuming a uniform configuration factor and holding x_i constant gives:

$$D_s = \frac{kqx}{C(1+k)^2} \sum_i s_i^2 = Z \sum_i s_i^2 \quad (20)$$

which reduces to:

$$D_s' = Zs^2 \quad (21)$$

for a uniform film of thickness s . For a distribution with an equal volume of film 0.1s and 10s in thickness:

$$D_s' = Z [5(0.1s)^2 + 0.05(10s)^2] = 5.05 Zs^2 \quad (22)$$

This distribution, resulting from the same volume of stationary phase present in the uniform thickness case, leads to a diffusivity value five times larger than that corresponding to a uniformly distributed film. Table VI-4 summarizes the effects of different thickness distributions on the diffusion coefficient D_s' compared to that expected from a uniform film of thickness s , D_s . All the geometries considered require the same volume of stationary phase per unit column volume and have an average thickness of s . It is evident from these examples that thick sections of the polymer film

Table VI-4
Effect of Thickness Distribution on Diffusivity
in the Polymer Phase

Thin Section Depth	Thick Section Depth	D_s^1/D_s
0.50s	2s	1.25
0.25s	4s	2.13
0.10s	10s	5.05
0.02s	50s	25.01
0.01s	100s	50.01

can exert a disproportionate influence and lead to a considerable increase in the polymer-probe diffusion coefficient when compared to the value calculated for an IGC experiment with an assumed ideal polymer layer of uniform thickness. An experimentally difficult determination of the variability of the polymer film in the IGC column would be necessary in order to correct the calculated diffusivity values for the actual polymer layer nonuniformity.

Experimental Details

The column used throughout the relative diffusivity study was prepared from glass support beads coated with a layer of the linear polyethylene (LPE) NBS SRM 1475 and packed into a 6.35 mm outer diameter stainless steel tube, as described in the previous annual report [1]. After installation in the gas chromatograph, a final drying and column conditioning step was carried out by purging with dry, deoxygenated helium followed by heating to 150 °C for a 24-hour period. The column used is identical to column B in the previous annual report and contains 5.71×10^{-3} mass fraction of LPE. It was chosen because of the relatively small contribution which gas phase mass transfer made to the overall peak spreading when compared to column A of that report.

Several alkane probes and an antioxidant were employed as diffusants and are listed in the table below [16]. All were dissolved in hexane (Burdick and Jackson Labs.) to give solutions of about 0.02, 0.2, and 2.0 percent by mass.

Table VI-5
Probes Employed in an IGC Study of Relative Diffusivity in LPE

Probe	Code	Source
n-octadecane	C ₁₈	Supelco
n-hexadecane	C ₁₆	Aldrich Chemical
2-methylpentadecane	2MeC ₁₅	Alfa Products
2,2,4,4,6,8,8-heptamethylnonane	7MeC ₉	Aldrich Chemical
n-tridecane	C ₁₃	American Petroleum Institute
1,3-di-tert-butyl-2'-hydroxy-5-methylbenzene	BHT	Polysciences

Approximately 1 μ l of each solution was injected into the flash vaporization injector of a gas chromatograph, maintained at 180 °C. The column oven was held at 150 °C \pm 0.3 °C throughout. The eluted peaks were monitored with a flame ionization detector (FID) held at 190 °C, whose signal at 10^{-10} A/mV sensitivity was fed to an electronic integrator connected to a strip chart recorder and to a high acquisition speed recorder-plotter. The former setup was employed to time the peak maxima to within 0.01 min and for manual measurements of the chart width at half maximum at chart speeds up to 1 cm/s. Typical results are shown in Figure VI-4.

The rapid collection capabilities of the high speed recorder-plotter allowed the FID output to be sampled at 10 ms intervals, facilitating analysis

of the marker and solvent peaks. A typical trace is shown in Figure VI-5 for five repeated determinations. Data collection was triggered upon injection of the methane and hexane vapor sample and the time to the methane peak maximum, assumed to have $k = 0$, was employed to determine t_M of equation 6 and to calculate the column average linear flow velocity, \bar{u} , from the expression:

$$\bar{u} = L/t_M \quad (23)$$

The quantity of interest, the column outlet velocity, was then calculated from \bar{u} as follows:

$$u_o = \frac{2\bar{u}}{3} \left[\frac{\left(p_i/p_o \right)^3 - 1}{\left(p_i/p_o \right)^2 - 1} \right]. \quad (24)$$

Column inlet and outlet pressures were monitored with calibrated pressure gauges of appropriate range. The outlet pressure was very near to atmospheric in all cases. The helium carrier gas volumetric flow rate was measured with a soap film flow meter, when necessary. The data were collected over a corrected flow rate range of about 1 to 5 ml/s.

The first derivative of a chromatographic peak with respect to time can be used to aid in determinations of the maximum height and peak widths indicated in Figure VI-1. At a point where the chromatogram is at a maximum, the first derivative will be zero while the second derivative is negative. Determination of the first derivative zero point is illustrated with curve B of Figure VI-6, the first derivative of curve A obtained by experiment. The inflection points of the chromatogram are also readily indicated by maxima or minima in the derivative curve. The magnitude of the derivative at this point gives the slope of a tangent to the peak. The ratio of the absolute values of the derivative maximum and minimum gives a measure of peak symmetry.

For the purposes of this study, a symmetric peak is defined as that having a front to rear tangent slope ratio of 1 ± 0.2 . The highly asymmetric peaks that result from excessive probe concentrations or adsorption, two of the many factors which can cause deviations from Henry's law in the chromatographic column, are not amenable to analysis by the methods outlined here for Gaussian-shaped peaks. The time period between maximum and minimum of the derivative curve can be measured easily and is equal to twice the standard deviation of the peak, as illustrated in Figures VI-1 and VI-6. The above procedure was used to collect information on the center position, standard deviation, and symmetry of the eluted diffusant peaks.

Relative Diffusivity Determinations

As noted earlier, the probes listed in Table VI-5 were studied in the same LPE-containing column. This allows the diffusivity of one probe in LPE to be compared to that of another, irrespective of the method of any calculation of the absolute diffusivity of the probes in the polymer. The experimental data acquired for tridecane in LPE is given in Table VI-6 and can be used to give an indication of typical values of the various parameters utilized in these determinations. The variables p_i , p_o , and t_M are used in the calculation of u_c by equations 23 and 24. The plate height is determined from $w_{1/2}$ and t_R with equation 1 or from w_i and t_R by the following expression:

$$H = \frac{L}{4} \left(\frac{w_i}{t_R} \right)^2 \quad (25)$$

When values of A and B are available, the plot suggested by equation 15 can be constructed from the data of Table VI-6. The previous annual report [1] contains a more detailed example of the calculation of these two constants from equations 3 and 4 than the abbreviated discussion given here. The values used in calculating \bar{H}/u_o are given in a footnote to Table VI-6.

Table VI-6
Experimental Data for an IGC Diffusivity Study of Tridecane in Linear Polyethylene at 150 °C

Pressure (Torr)	Time to Maximum (s)		Peak Width (s)	Outlet Velocity	Plate Height	$\frac{\bar{H}}{H} / u_0, (ms)^*$	Ordinate	Abscissa
	P_i	P_o	t_M	t_R	$w_{\frac{1}{2}}$	w_i	$u_0 (\text{cm/s})$	
3160	745.9	4.23	54.6	8.68	7.34	108.2	0.705	6.31
2590	745.0	5.29	69.0	9.79	8.34	72.2	0.562	7.44
2103	743.4	6.94	90.6	11.10	9.45	46.0	0.419	8.49
1827	744.6	8.43	111.0	12.15	10.29	33.6	0.335	8.95
1577	745.2	10.78	141.0	13.73	11.68	23.3	0.265	9.56

*The following values were used in the calculation of $\frac{\bar{H}}{H} / u_0$: $L = 154.9 \text{ cm}$, $A = 0.0165 \text{ cm}$, $B = 0.585 \text{ cm}^2/\text{s}$

Plots of \bar{H}'/u_0 against the $2p_0/(p_i + p_0)$ term are shown for five diffusants in Figure VI-7. The slopes of these plots provide a value for the C constant needed in a diffusivity determination. These C values are listed in Table VI-7, accompanied by the corresponding value of k for each

Table VI-7

Relative Diffusivity Results for Several Alkanes and BHT in LPE at 150 °C

Probe	C (ms) $\pm \sigma$ (%)	k $\pm \sigma$ (%)	D _{Probe}	D _{Probe} /D _{C₁₈}	D ($10^7 \cdot \text{cm}^2/\text{s}$)
C ₁₈	21.9 \pm 7	119 \pm 3	0.38 Y	1	2.4
C ₁₆	7.8 \pm 14	60.9 \pm 1	2.05 Y	5.4	13
2MeC ₁₅	8.7 \pm 12	49.0 \pm 1	2.25 Y	6.0	14
7MeC ₉	18 \pm 11	12.8 \pm 1	3.74 Y	9.9	24
C ₁₃	12 \pm 9	12.1 \pm 1	5.74 Y	15.2	36
BHT	9.9 \pm 11	32.4 \pm 2	2.94 Y	7.8	19

probe as calculated from the data of Table VI-6 by equation 6. This information can then be used to determine a value for the relative diffusivity by the following procedure. Equation 17 can be rearranged to collect all the variables used only to characterize the polymer packing into one term and solved for the diffusivity. The following expression results when this is done:

$$D_s = \frac{k}{C(1+k)^2} \left(\frac{r^4 m_s \rho}{168.75 \rho_s} \right)^{\frac{1}{2}} \quad (26)$$

The first term in the above equation contains the variables which characterize the interactions between the probe and polymer while the square root term consists of variables which describe polymer layer and support

particle characteristics only. These latter variables are constant throughout all determinations with this particular column and this collection of parameters can be abbreviated as the constant Y. The following version of equation 26 is consequently obtained:

$$D_s = \frac{k}{C(1 + k)^2} \cdot Y \quad (27)$$

This equation was used to determine the probe diffusivity values given in the fourth column of Table VI-7. If one of the probes is selected as a reference, in this instance octadecane, the diffusivity of a probe in the polymer relative to the diffusivity of the reference probe in that polymer can be obtained from a ratio of the respective results from equation 27. These values are given in the fifth column of Table VI-7.

An alternate determination of the relative diffusivity might be based on equation 7. In this instance the following expression would be obtained for D_s :

$$D_s = \frac{k}{C(1 + k)^2} \cdot \frac{2s^2}{3} = \frac{k}{C(1 + k)^2} \cdot Y' \quad (28)$$

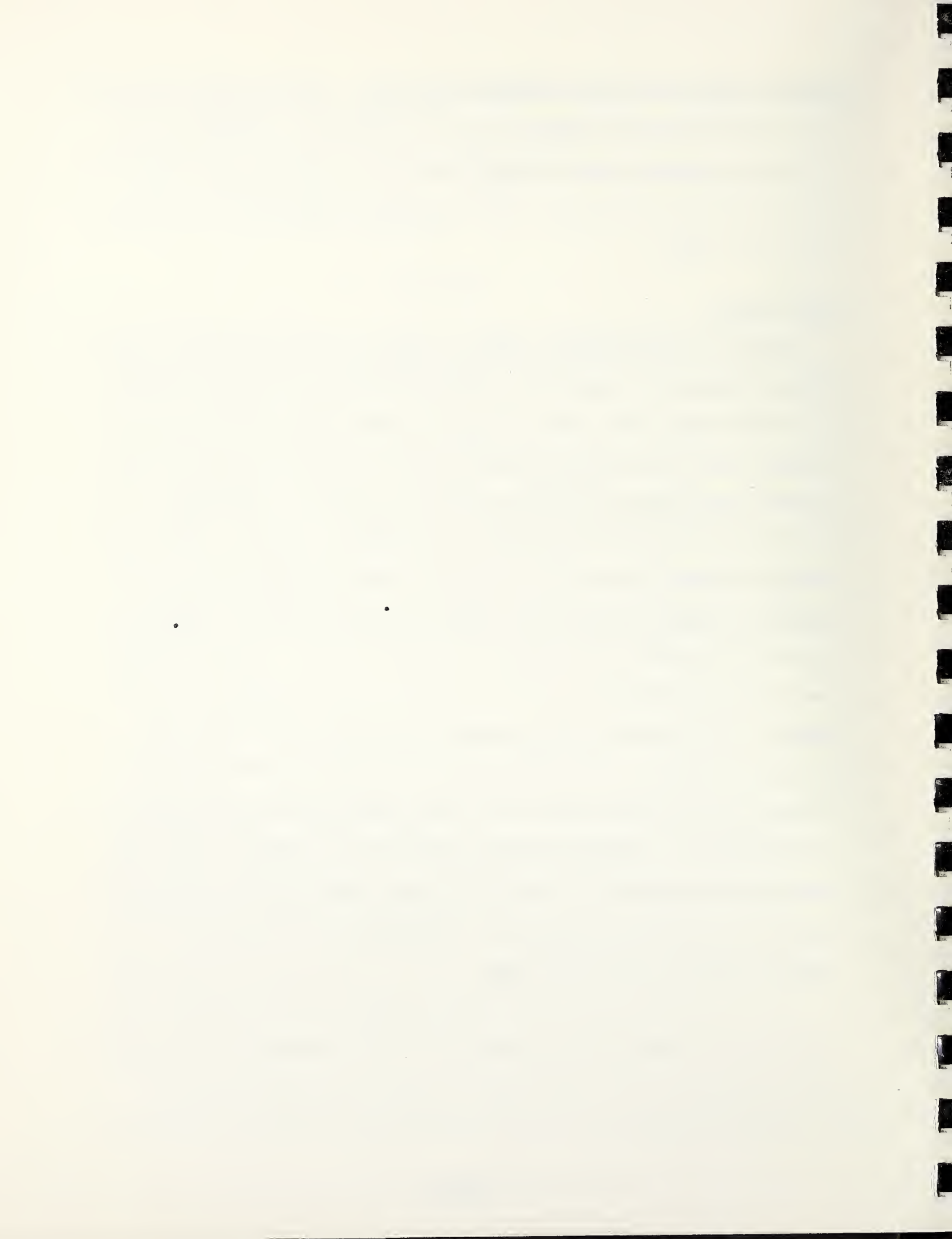
Equations 27 and 28 are identical except for the constants Y and Y', which are derived from different collections of parameters used to describe the distribution of the polymer phase on the support. The numerical front-factor for Y' is identical to that given for Y in column four of Table VI-7 since it depends only on the values of C and k, which are the same for either method. The relative diffusivity results would also be identical as the Y or Y' constant is eliminated by the ratioing method used to obtain this quantity.

An absolute diffusion constant for each probe in the polymer can readily be obtained from the relative diffusivity values if an absolute diffusion coeffi-

cient for any of the probes in that polymer exists. Taking the previously cited value in Table VI-3 for column B of $2.4 \times 10^{-7} \text{ cm}^2/\text{s}$ for octadecane in LPE at 150 °C as accurate, the absolute diffusivities in column six of Table VI-7 were obtained by multiplying the relative diffusion coefficients of column five by this factor.

Future Results

Results on lower molecular weight alkanes, principally isomers of octane and normal decane and tridecane, at temperatures below the LPE melting point will be given in a later report. They will allow a more rigorous evaluation of the IGC relative diffusivity technique as many more absolute diffusivity determinations have been conducted with these compounds. Independent values for the absolute diffusion coefficient of tridecane in LPE will then be available at two temperatures, one below, and one above the polymer melting point. By using an appropriate activation energy to extrapolate one of the results and making allowances for the expected effects of polymer crystallinity on the diffusion coefficient, a direct comparison of the two results will be possible. Unfortunately, it is not possible to study the alkanes from octane to octadecane at a single temperature due to the great differences in volatility among these compounds. This necessitates the temperature extrapolation for a common probe described above as a means of connecting the two bodies of data collected at different temperatures and allowing for a critical evaluation of the method. Additional results for BHT at lower temperatures will also be provided.



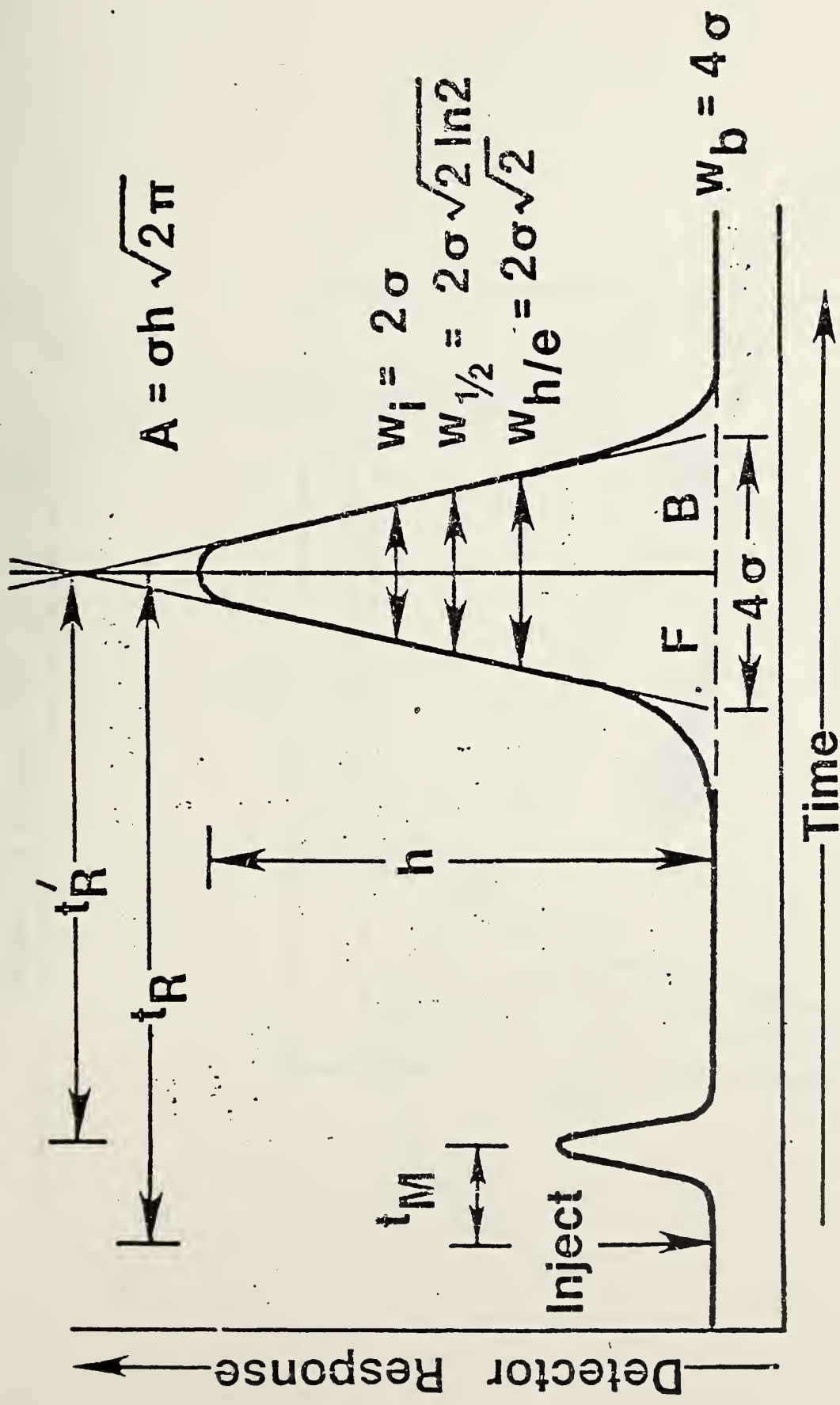


Figure VI-1: Elution chromatogram schematic showing a marker peak at t_M and a Gaussian probe peak at t_R . Various methods of determining the standard deviation σ are indicated.

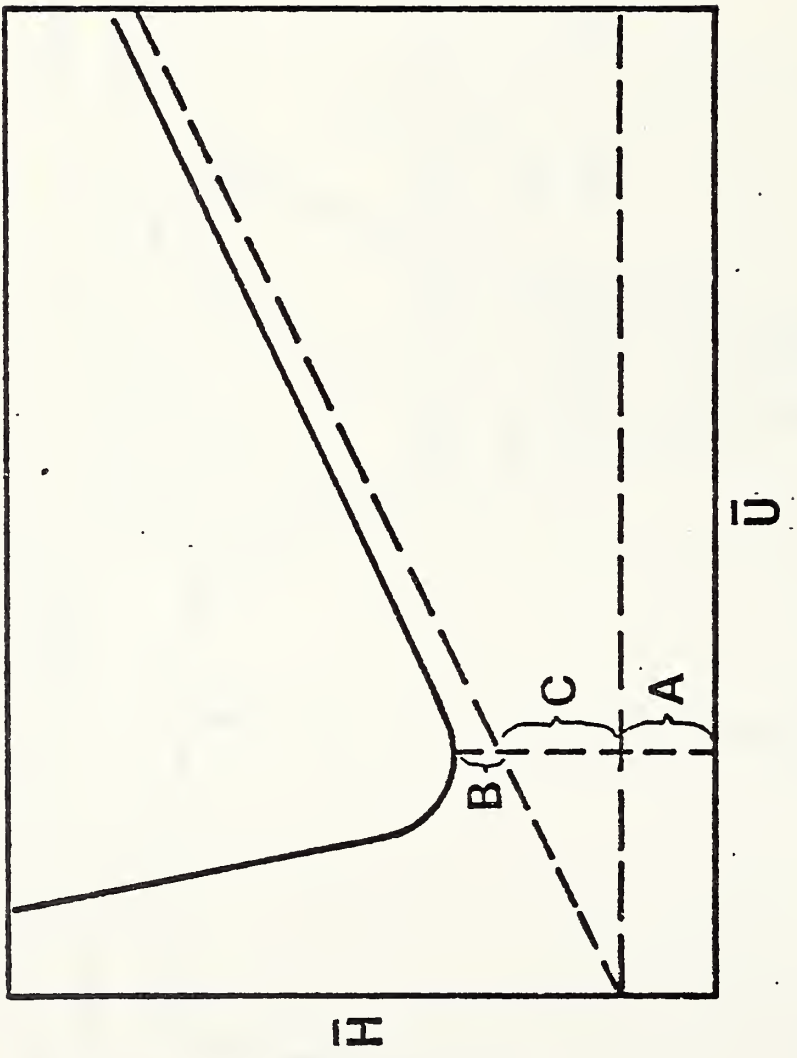


Figure VI-2: van Deemter plot of the height equivalent to a theoretical plate against the average carrier gas flow velocity.

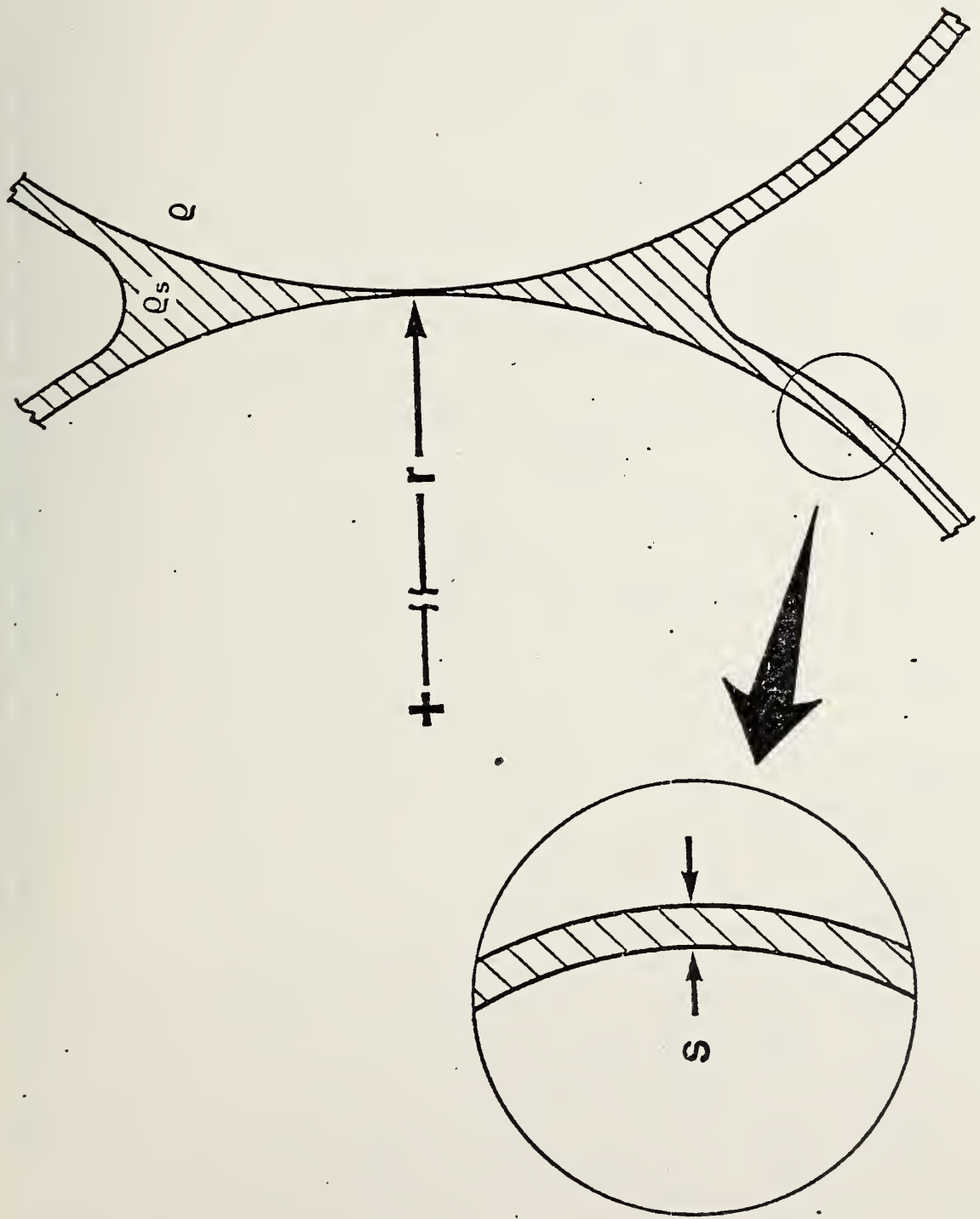


Figure VI-3: Illustration of the possible polymer phase geometries within an IGC column containing a glass bead support.

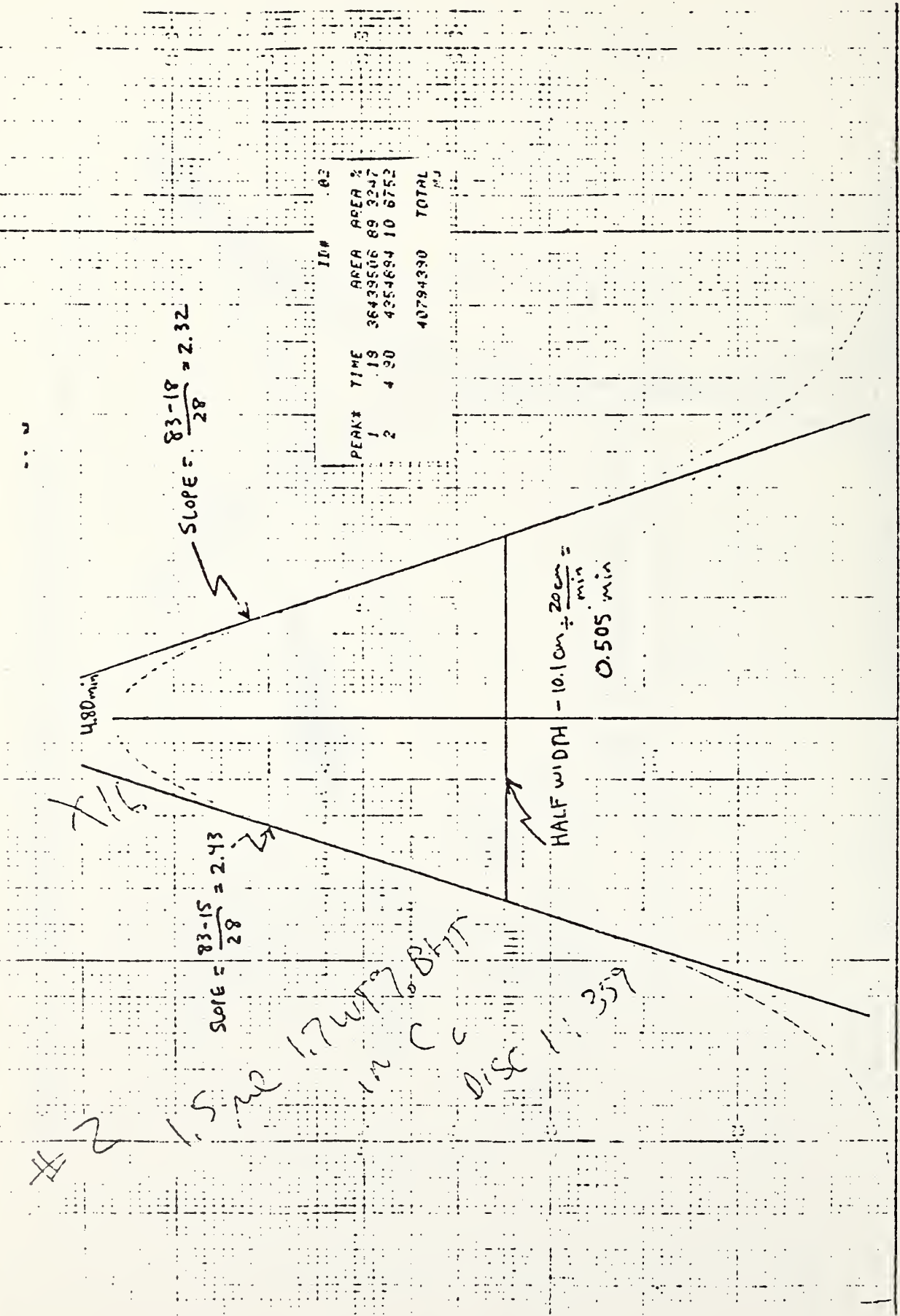


Figure VI-4: Typical strip chart recorder output of a probe peak showing the methods for determining the width at half maximum and tangent slopes.

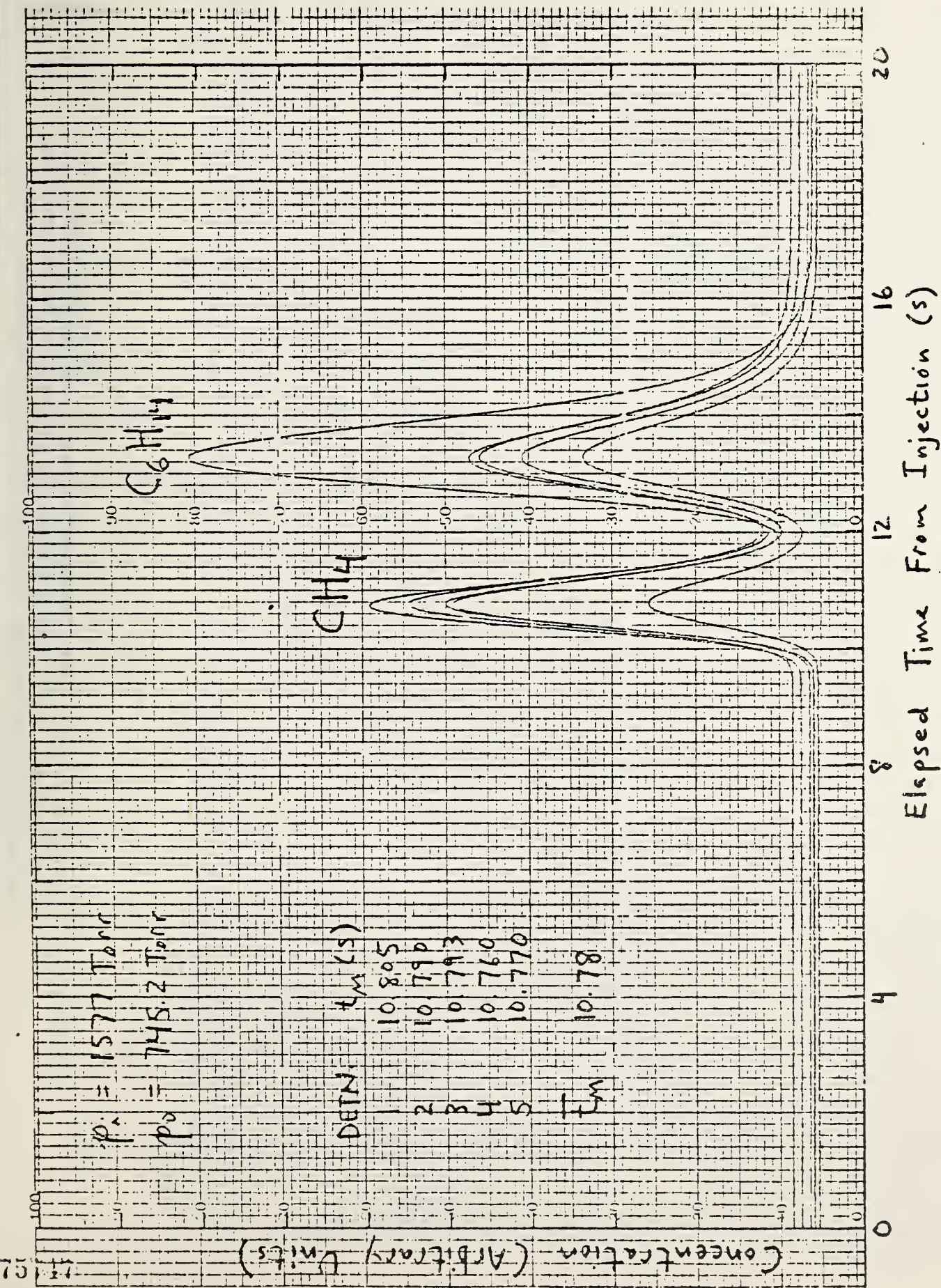


Figure VI-5: High speed chart record of five determinations of t_M with methane as the marker. A hexane solvent peak is also present.

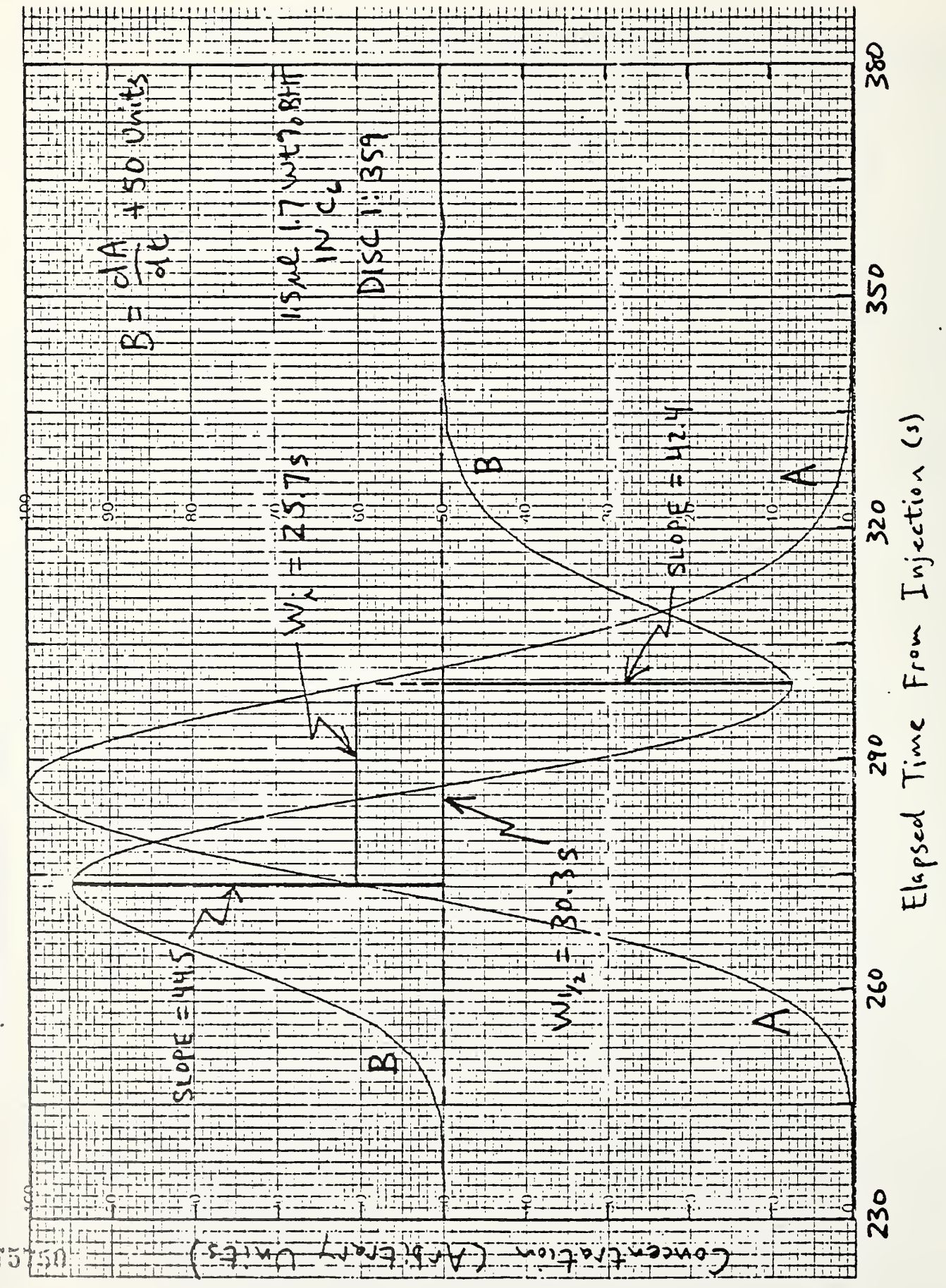


Figure VI-6: Typical high speed chart record of a probe peak (curve A) and first time derivative (curve B) showing the methods for determining the width at half maximum, width at the inflection points, and tangent slopes.

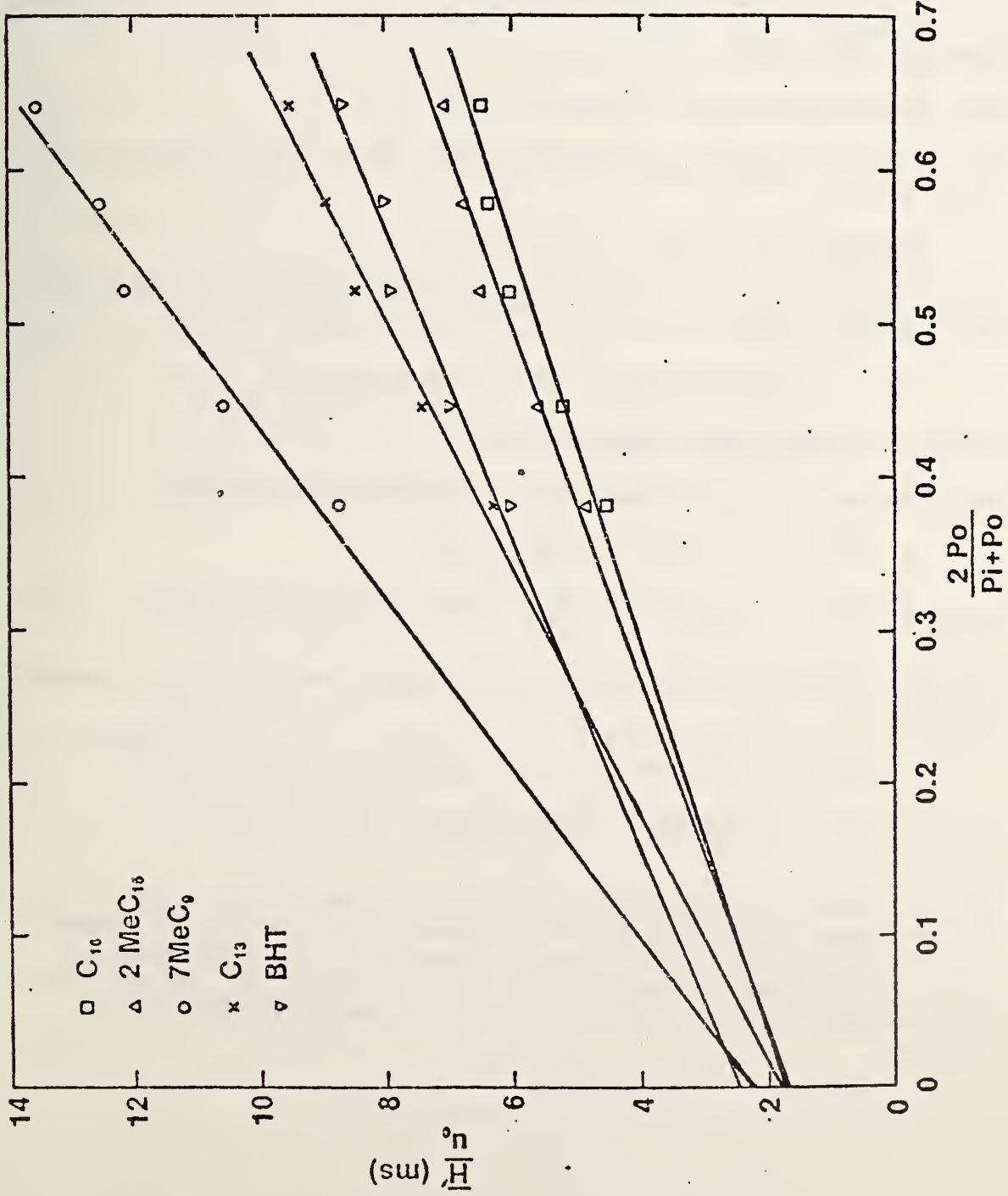


Figure VI-7: Plot of \bar{H}/u_0 vs $2P_o/(P_i + P_o)$ suggested by equation 15 for five diffusants in LPE at 150 °C.

References and Notes

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16. Certain commercial equipment, instruments, or materials are identified in this paper in order to adequately specify the experimental procedure. Such identification does not imply recommendation or endorsement by the National Bureau of Standards, nor does it imply that the materials or equipment identified are necessarily the best available for the purpose.

VII. A Simplified Migration Decision Tree

One of the most important parameters in indirect food additive regulation is the maximum allowable concentration or a critical concentration of the species of concern in food, C^* . This value is based upon toxicological studies and from the fractional amount of total food consumed, that would be in contact with the packaging material of concern.

Acceptable migration would be the case in which an upper bound level of migration for the maximum time and temperature for storage and usage, based upon either actual tests or sound estimations, is less than the critical or threshold level.

To this end, we have exemplified and simplified our decision tree described earlier (NBSIR 79-1779) as a flow chart shown in Figure VII-1.

1. Load Limited Migration

In the ultimate case, all the migrant of concern moves from the packaging material into food. The concentration of the migrant in food is:

$$C = \frac{V_p C_0}{V_s}$$

where C_0 is the original concentration of the migrant in the packaging material, V_p and V_s are the volumes of the packaging material and food or solvent respectively. Migration is acceptable if this concentration is less than the critical concentration, C^* .

In general, 10 g food is considered in contact with 1 in² (6.45 cm²) of the packaging material (CRF Title 21, Part 21, Part 177). In other words, by approximating the density of foods as 1 gcm⁻³, the ratio of the volumes of the packaging material to food V_p/V_s is 0.645 & where & is the thickness of the packaging material in cm. Commonly this ratio is less than 1:40 or the thickness of the packaging material is less than 0.04 cm.

Therefore, the migration level is acceptable if the following conditions are met:

$$C_0 < \frac{V_s}{V_p} C^* \quad \text{for all cases,}$$

or

$$C_0 < \frac{1.53}{\ell} C^* \quad \text{if no less than 10 g of food is in contact with } 6.45 \text{ cm}^2 \text{ of packaging material,}$$

or

$$C_0 < 40C^* . \quad \text{If } V_p \text{ is less than } 1/40 \text{ of } V_s .$$

2. Solubility Limited Migration

For the case in which the loading of the migrant of concern is greater than the above limit, the saturation solubility C_{sat} of the migrant in the particular type of food or food simulant is examined in order to find out whether the saturation solubility is less than the critical concentration. Special attention should be given to cases where surfactant or emulsification action may tend to increase the actual migrant concentration in food beyond the saturation concentration. The migration level is acceptable if

$$C_{sat} < C^* .$$

3. Partition Limited Migration

If both the loading and the solubility exceed the limiting threshold, then the partitioning of the migrant between the packaging material and the food or the simulant should be examined. The partition coefficient k is defined as the quotient of the activities, a , of the migrant in the food and in the polymer at equilibrium. It may also be approximated by the ratio of the solubilities in the solvent and in the polymer,

$$k = \frac{1}{K} = \frac{a_s}{a_p} \approx \frac{C_{sat}}{C_{p,sat}}$$

By defining α as the ratio of the amount of migrant in the solvent, M_∞ , versus that in the polymer, $M_{p\infty}$, at equilibrium

$$\alpha = \frac{M_\infty}{M_{p\infty}} = \frac{V_s}{V_p} k = \frac{V_s C_\infty}{V_p C_{p\infty}},$$

then the final fraction of original amount of additive migrated,

$$\frac{M_\infty}{M_0} = \frac{\alpha}{1 + \alpha}$$

where M_0 is the original loading of migrant in the polymer. For the final concentration in food or solvent to be less than the threshold,

$$C_\infty = \frac{M_\infty}{V_s} < C^*,$$

or
$$\frac{\alpha}{1 + \alpha} \frac{V_p}{V_s} C_0 < C^*,$$

it requires that
$$C_0 < \left(\frac{V_s}{V_p} + K \right) C^*.$$

Thus step 1 is just a special case of step 3.

In the case that the partition coefficient or the saturation concentration of the migrant in the polymer is unknown or difficult to determine, the partition coefficient may be assigned a value as in our previous reports (NBSIR 79-1598 and 79-1779)

$$\ln k = \ln C_{\text{sat}} + X_p^\infty + 1 = \ln C_{\text{sat}} + 2$$

In general, X_p^∞ is in the range of 0 to 2. Thus
 $k < 20 C_{\text{sat}}$.

This assigns a solubility of $C_{p,\text{sat}}$ of > 0.05 . Therefore the requirement is simplified as,

$$C_0 < (K + 40) C^*$$

or $C_0 < \frac{1}{20C_{\text{sat}}} + 40 C^*$

However, this simplification is applicable only when the saturated solubility in the food or solvent is relatively low and the solubility in the polymer relatively high.

4. Diffusion Limited Migration

When the equilibrium properties such as the additive loading in the polymer, the solubility of the additive in the solvent, and the partitioning of the additive between the polymer and solvent all yield too high a final concentration of the additive in the food or the solvent beyond the threshold concentration C^* , then it is necessary to find out kinetically whether, under the particular storage condition and in the limited storage time, the amount of the additive migrated would produce an additive concentration in the solvent less than that of the threshold limit. The kinetics of migration depends upon additive, polymer, and solvent combinations, as well as effects of temperature and stirring. The combination of additive, polymer, and solvent or food is the most important factor influencing the diffusion coefficient. This diffusion coefficient is also dependent upon temperature, with an activation

energy in the order of 10-20 kcal/mol for polymers in the rubbery state. A further relatively minor influence in comparison to the above factors may be observed due to the changes in the degrees of crystallinity and molecular weights of the polymer. Stirring speeds up the approach of an equilibrium condition.

If the migration is kinetically limited, i.e. much less than half of additive is migrated, then we are only interested in the short time behavior; otherwise, the estimation of total migration from step 1 would be sufficient.

For a single-sided extraction, at t when $M_t/M_\infty < 0.5$, we may apply the simple equation

$$\frac{M_t}{M_\infty} = 2 \left(\frac{Dt}{\pi} \right)^{1/2} \frac{1}{l}$$

In other words

$$\frac{M_t}{A} = 2 \left(\frac{Dt}{\pi} \right)^{1/2} C_0,$$

or

$$C_t = 2 \left(\frac{Dt}{\pi} \right)^{1/2} \frac{A}{V_s} C_0,$$

where l is the thickness of the film and A is the area of contact. For an upper limit, D is assigned the value of the diffusion coefficient measured on the polymer swollen by the solvent concerned. In the upper bound case, all migrants are assumed to be able to migrate from the polymer into solvent, i.e., $M_\infty = M_0 = V_p C_0$. If partitioning exists, M_∞ should be reduced from M_0 according to that described in step 3.

By assuming a storage time, t , of 1 year (3.15×10^7 s) and a container thickness of 0.04 cm (calculated from a food to container contact area of 10 g/in² and a container to food weight ratio of 1:40), one may obtain

$$M_t/M_\infty \approx 1.6 \times 10^5 D^{1/2},$$

or

$$M_t/A \approx 6300 D^{1/2} C_0,$$

or

$$C_t \cong 4100 D^{1/2} C_0.$$

For $M_t/M_\infty = 0.5$, a diffusion coefficient of $D = 10^{-11} \text{ cm}^2 \text{s}^{-1}$ is required to meet the threshold limit. Therefore, with the above mentioned storage conditions, only in the cases where D is less $10^{-11} \text{ cm}^2 \text{s}^{-1}$ will diffusion limited migration be important enough to be considered. Otherwise equilibrium conditions should be used for estimation. For thinner packaging materials, D should be even less or at less than $6.2 \times 10^{-9} \text{ cm}^2 \text{s}^{-1}$ in order for the kinetic criteria for migration to apply.

For polyolefins, the glass transition temperature is lower than -20°C . Hence, the semicrystalline solid polymers are, in general, utilized in the condition of mixed rubbery (amorphous portion) and crystalline states. It is usually assumed that the additive migration occurs in the amorphous region. The diffusion coefficients of migrant for the rubbery state are generally many orders of magnitude higher than those for the glassy state. For oligomers up to $n\text{-C}_{32}\text{H}_{66}$ and anti-oxidant molecules such as BHT, the apparent diffusion coefficients into either corn oil, triglycerides or ethanol are near or greater than $10^{-11} \text{ cm}^2 \text{s}^{-1}$ above 30°C . The influence of solvent, temperature, and crystallinity on the diffusion coefficients of BHT in polyethylene have been summarized earlier in this report. Therefore, for polyolefins, the diffusion limited case exists for these migrants and solvents only at a storage temperature lower than ambient, or at a much shorter storage time (a 10-fold reduction in the amount migrated requires a 100-fold reduction in time), or for a very thick container. The diffusion limited case may also exist for extraction with poor solvents for the migrant. However, in the case with poor solvents, the solubility limit or partition limit requirements may often suffice.

Fig. VII-1. A Simplified Additive Migration Decision Flow Chart

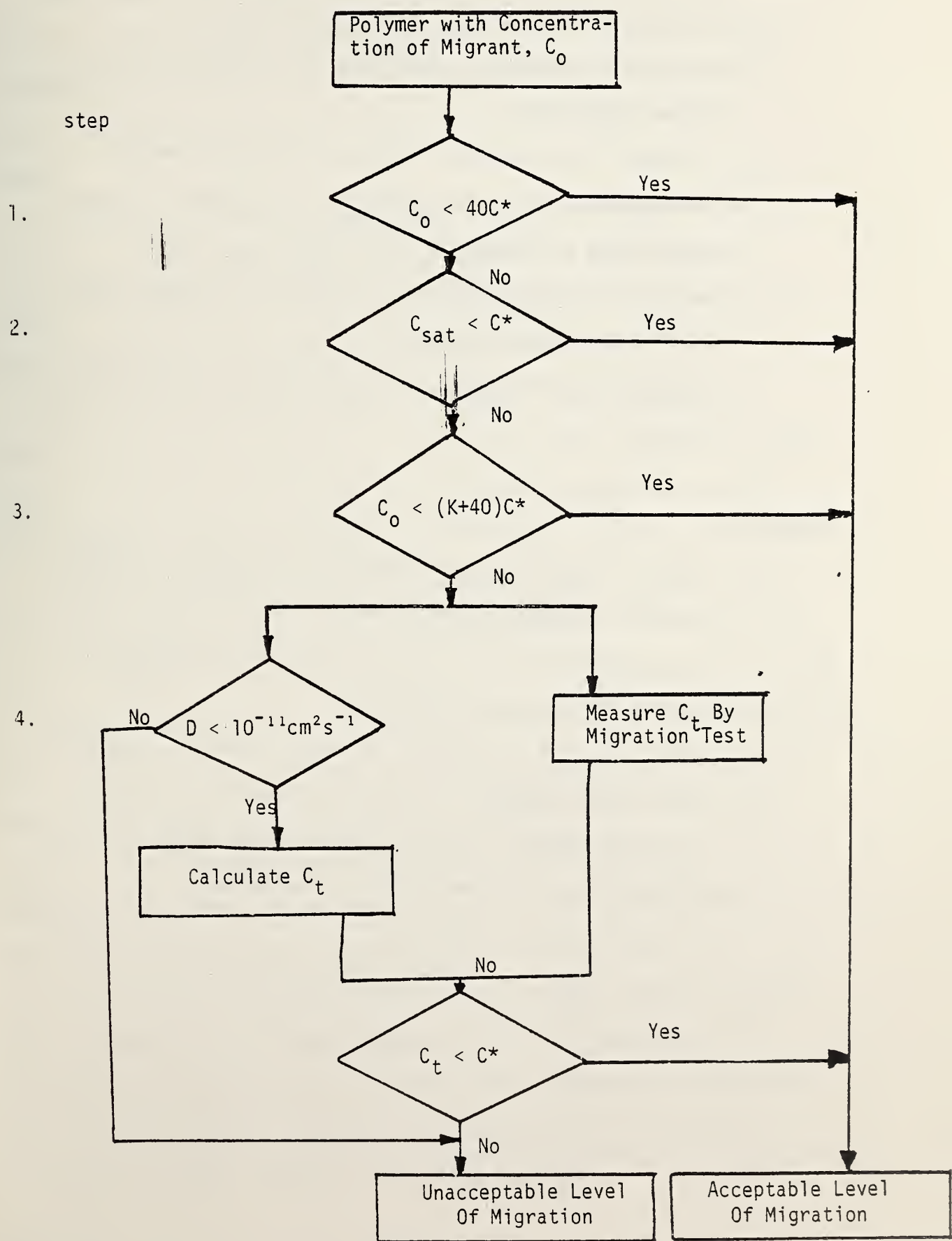


Figure VII-1. (Continued)

Glossary:

C_0 - Original concentration of migrant in polymer.

C^* - Critical or acceptable concentration of migrant in food or solvent of concern.

C_{sat} - Solubility of migrant in food or solvent of concern.

K - Distribution or partition coefficient of migrant in polymer and in food or solvent at equilibrium, C_{poly}/C_{solv} .

D - Apparent diffusion coefficient of migrant in polymer chain by food or solvent of concern.

C_t - Concentration of migrant in food or solvent after contact time t.

Notes:

1. This flow chart is based upon the assumption of a volume ratio of packaging material/food of 1:40 or 10 g of food are in contact with 1 in² or 6.45 cm² of packing material. Consult text for other conditions.
2. Along the decision or branching points, move to the next step if the parameter in the particular step is unknown or not determined, the particular step may be skipped and the criteria of the next step will be tested.

VIII. Simulating Solvents

Within the limited temperature range of food utilization and storage, solvent is the single most important parameter effecting both the amount of migrant extractable and the diffusion coefficient of the migrant, acting through its solubilities in and its swelling action upon the polymer.

By correlating diffusion coefficients in other solvents to that in one of the solvents, e.g. ethanol, while keeping all other parameters constant, it is possible to reduce the large spread of diffusion coefficients of the order of 10^8 to within a factor of 5 or less.

For partitioning or low solubility solvents, the correlations depend not only on the solvent but also on the migrant and/or the polymer. However, for solvents having moderate solubilities toward the migrant, the correlation seems to be independent of migrant, polymer or temperature.

The most significant correlations are those found for pure triglycerides and ethanol as food oil simulants and that for the accelerating solvent, n-heptane. As shown in Figure VIII-I, the diffusion coefficients of migrants (n-octadecane, n-dotriacontane or BHT) moving from polyolefins (LPE, BPE, PP) or E-VA copolymers into either tributyrin, trioctanoin or synthetic triglyceride mixture HB307 at either 30 or 60 °C are almost the same as the corresponding diffusion coefficient in corn oil. The diffusion coefficients in the pure triglycerides are slightly higher than that in corn oil with a ratio of $D(\text{pure triglycerides})/D(\text{corn oil})$ less than 1.5 over most of the range studied. Ethanol seems to behave almost identically to the lower members of pure triglycerides in its extraction behavior as shown in Figure VIII-2. However the correlation for the data of ethanol extractions shows more scatter than that for triglycerides within a factor of less than 3 or less.

From the above observations, either ethanol or low members of pure triglycerides may be used successfully for most occasions as food oil simulants, regardless of dissimilarities in viscosities, solvation power for the migrants, and swelling power for the polymers.

The accelerating action of n-heptane over that of food oil has long been recognized. The amount extractable by food oil was sometimes considered to be 1/5 of that by n-heptane. However we found that no constant factor will describe this accelerating action. Beside the differences in solubilities and in partition coefficients, the accelerating power of n-heptane is clearly demonstrated in Figure VIII-3, where diffusion coefficients in n-heptane and in triglycerides (including corn oil) are plotted against those in ethanol for identical test samples and conditions. Regardless of polymer, migrant and temperature, the accelerating power is greater at low diffusion coefficients than that at high diffusion coefficients. While, at $D(\text{ethanol})$ around $10^{-7} \text{ cm}^2 \text{s}^{-1}$, accelerating power is greater at low diffusion coefficients than that at 1000 at $D(\text{ethanol})$ around 10^{-11} or $10^{-12} \text{ cm}^2 \text{s}^{-1}$.

The accelerating action of n-heptane compared to that of ethanol (and that of triglycerides), coupled with the differences in solubilities of oligomers of polyolefins, has dramatic effects on the total extractables of polyolefins. We, therefore, subjected large quantities (50 g) of the raw polymer stocks of SRM 1475 (LPE) and SRM 1476 (LPE) to extraction by 500 ml of ethanol n-heptane at 70 °C for 160 days with occasional shaking. By assigning a minimum diffusion coefficient of about $10^{-9} \text{ cm}^2 \text{s}^{-1}$ for $n\text{-C}_{32}\text{H}_{66}$ and a pellet radius of 0.1 cm, the effective time $T = Dt/\ell^2$ is 1.4. Thus according to the generalized solution for the diffusion equation, the extraction should be at least 95% complete. The results of the 160-day extractions are listed in

Table VIII-I. For each polymer, about 6 to 8 times as much of polymer fraction was extracted by n-heptane than by ethanol.

The extracts were then subjected to gel permeation or size exclusion chromatographic analysis by Roger C. Snyder of FDA to find out their molecular weight distributions, as summarized in Table VIII-2 and shown in Figure VIII-4 for linear polyethylene SRM 1475 extracts, respectively. The molecular weight scales were calibrated by means of analyzing five NBS polyethylene standard reference materials, three n-alkanes (C_{10} , C_{20} , C_{44}) and test mixtures containing known amounts of n-alkanes from C_6 to C_{44} . The molecular weight scale for branched polyethylene is only approximately correct, however it should be sufficient for comparison purposes.

The extracts from the linear polyethylene show rather normal symmetrical distributions with peaks of molecular weight centering at about 300 for ethanol and at about 800 for n-heptane extracts. Such results are expected from the practice of molecular weight fractionation by solvents of different solubility power. On the low molecular weight side, e.g., $n-C_{18}H_{38}$, there is a large enough solubility or even total miscibility of the oligomer fraction in both solvents, such that almost all of the low molecular weight fraction will be dissolved by the extraction processes, cf. labels A in Figures VIII-4 and VIII-5. As molecular weight becomes higher, e.g., $n-C_{32}H_{66}$, the solubility in the solvent, especially in ethanol, decreases which causes a partitioning of the particular component between the polymer and the extracting solvent to occur, cf. labels B in Figures VIII-4 and VIII-5. The partitioning of $n-C_{32}H_{66}$ has been confirmed by extraction experiments with ethanol. The diffusion coefficients for larger molecules also drop significantly. Thus the combination of the effects of solubility, partitioning and diffusion created the shape of the molecular weight distribution curve beyond the peak molecular weight extracted.

A similar comparison for branched polyethylene is seen in Figure VIII-5. The skewed curves are most likely due to the incorrectness of the size versus molecular weight calibration which was based on n-paraffins and linear polyethylene fractions. The peak of apparent molecular weight extracted is about 350 for ethanol and about 3200 for n-heptane extracts.

Therefore, n-heptane can not only accelerate the diffusion process but also remove high molecular species of oligomers that may be hardly present in ethanol or triglyceride extracts.

Table VIII-1

Extraction of Polyethylenes

by solvents at 70°C. for 160 days

<u>Solvent</u>	<u>Fraction Extracted</u>	
	LPE SRM 1475	BPE SRM 1476
Ethanol	0.0008	0.0023
n-Heptane	0.0047	0.018

Table VIII-2
Molecular Weight Distributions of Ethanol and
n-Heptane Extractions of Polyethylenes

<u>Solvent</u>	$M_n : M_w$	
	LPE SRM 1475	BPE SRM 1476
Ethanol	250 : 310	310 : 410
n-Heptane	560 : 1020	620 : 3000

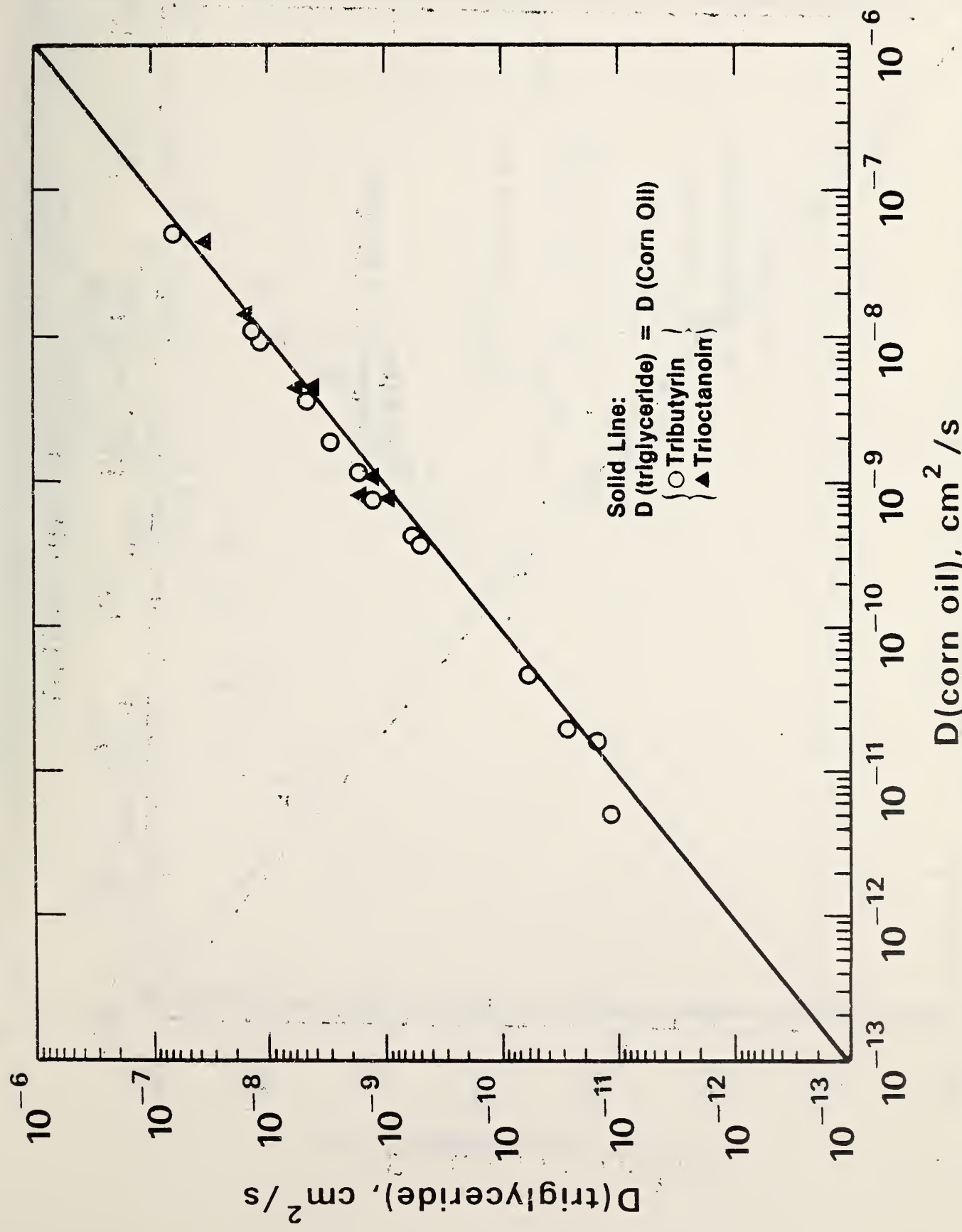


Figure VIII-1 Comparison of Diffusion Coefficients in Triglycerides and in Corn Oil

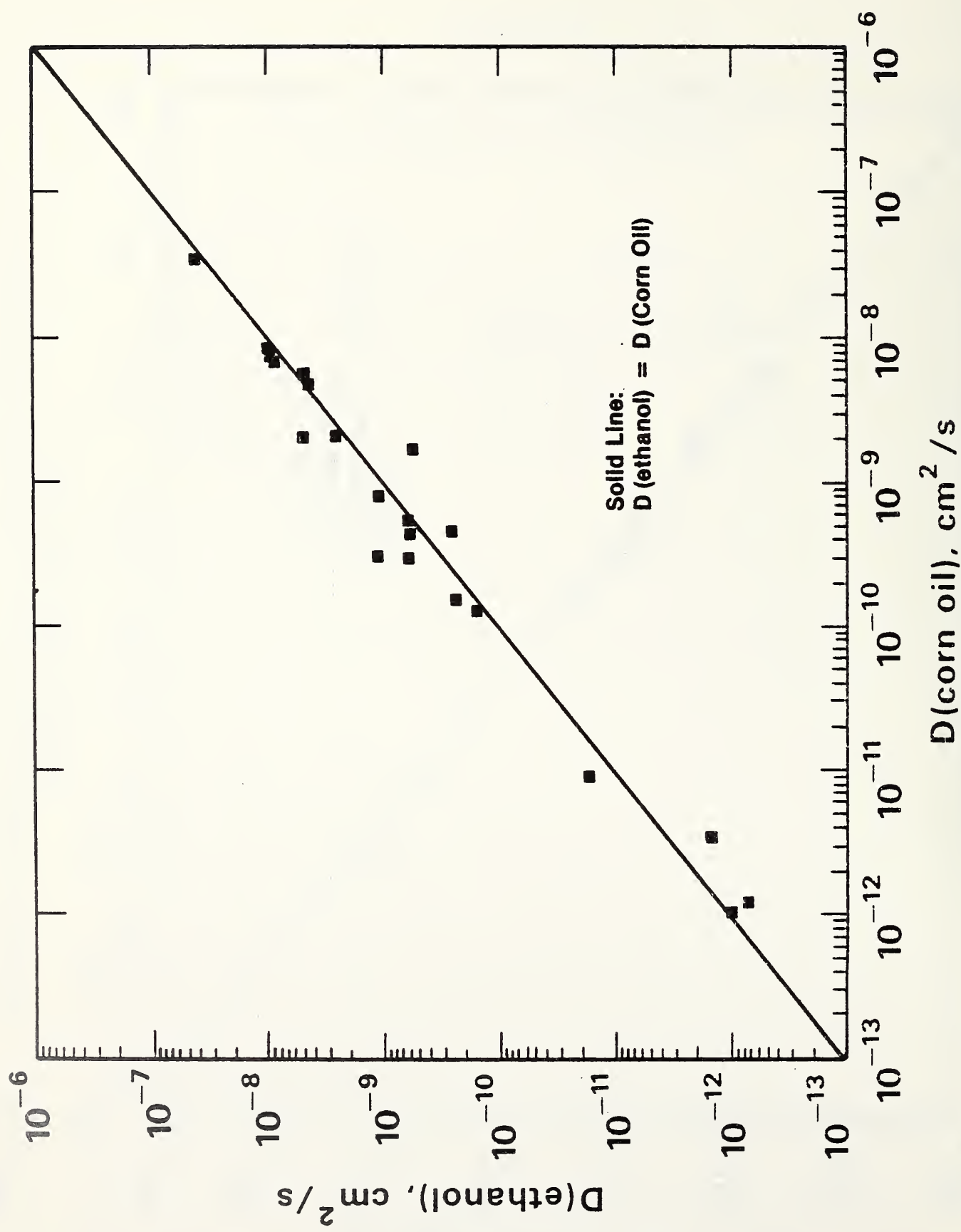


Figure VIII-2 Comparison of Diffusion Coefficients in Ethanol and in Corn Oil

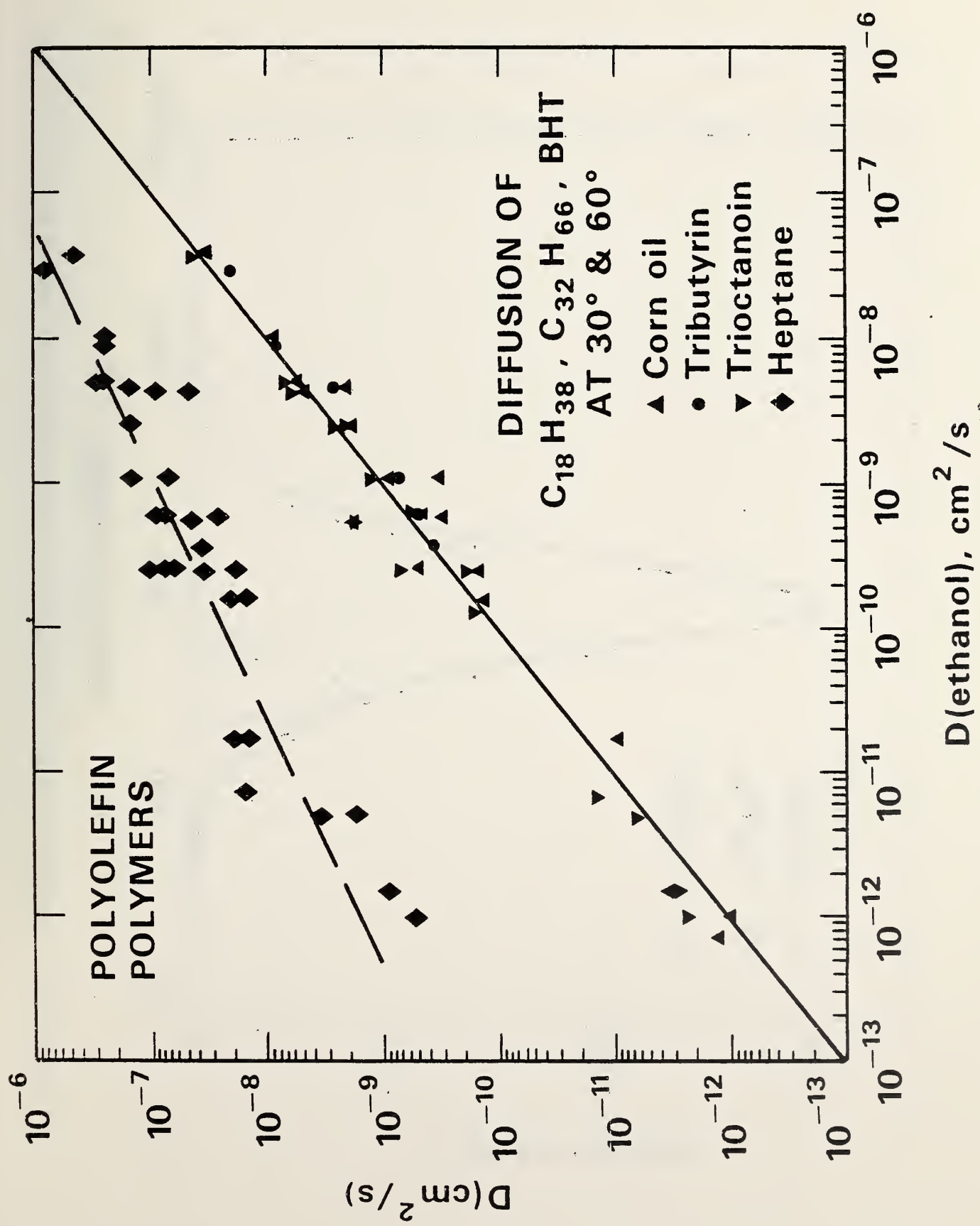


Figure VIII-3 Effect of Accelerating Solvent

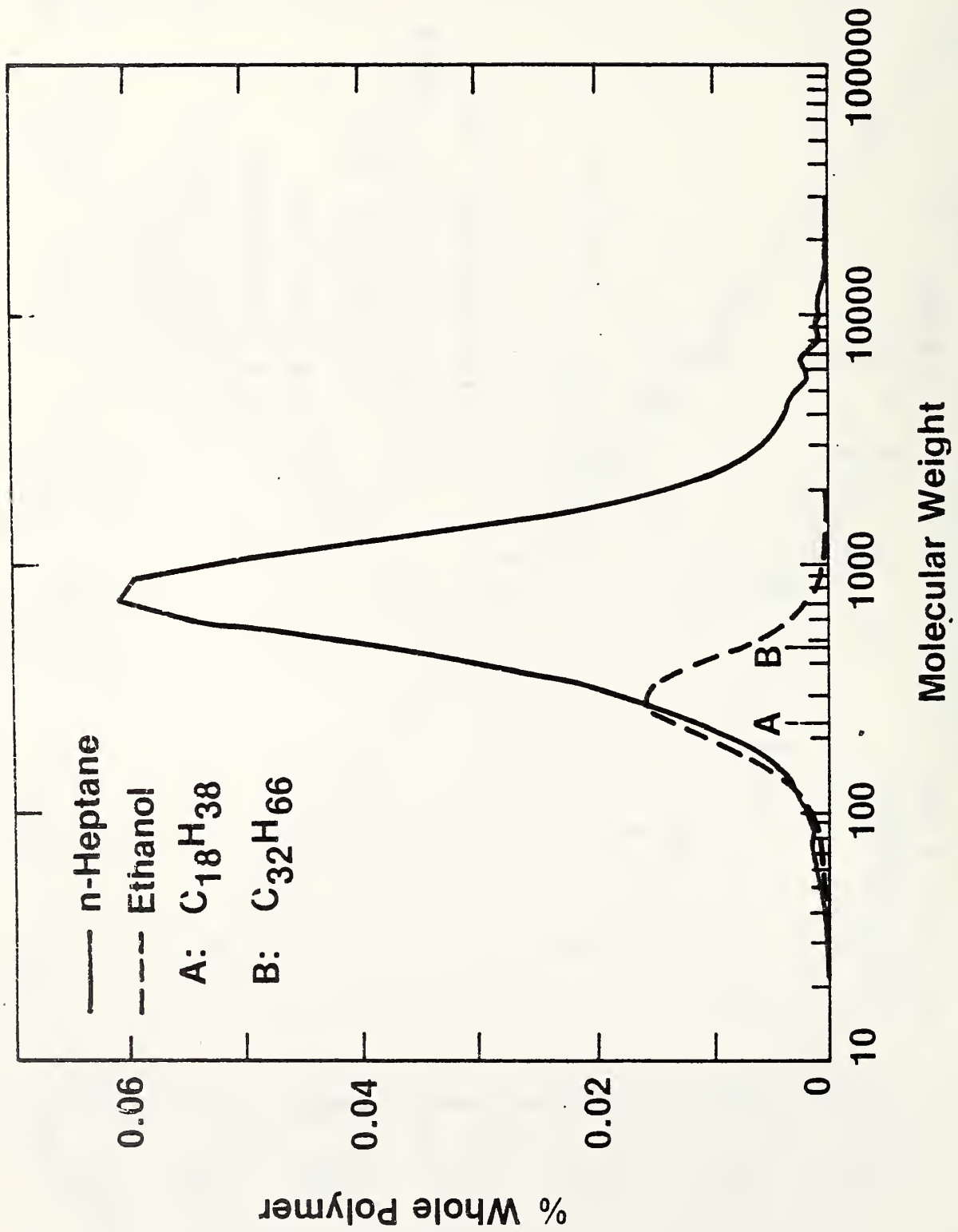


Figure VIII-4 Molecular Weight Distributions of Ethanol and n-Heptane Extracts of LPE

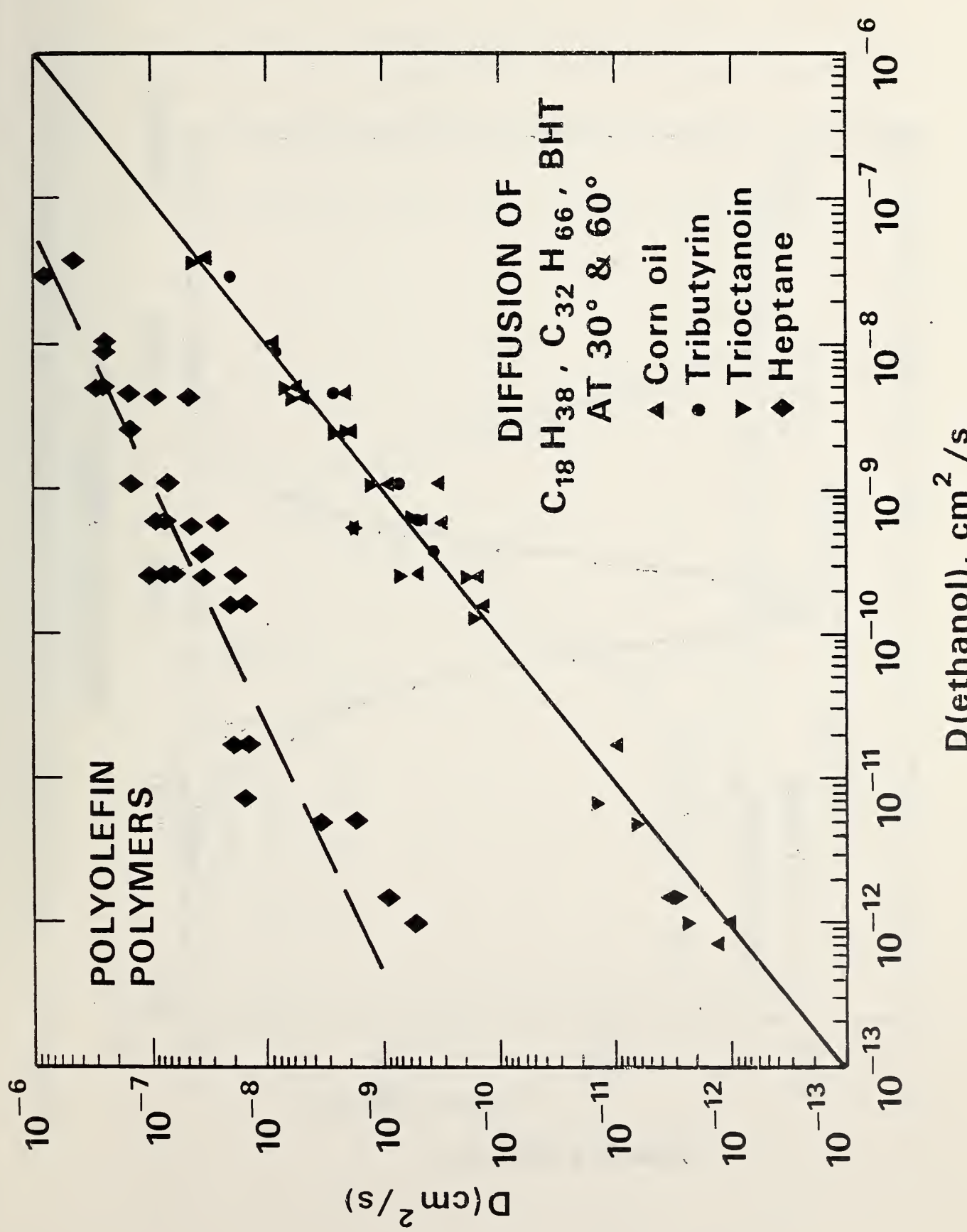


Figure VIII-3 Effect of Accelerating Solvent

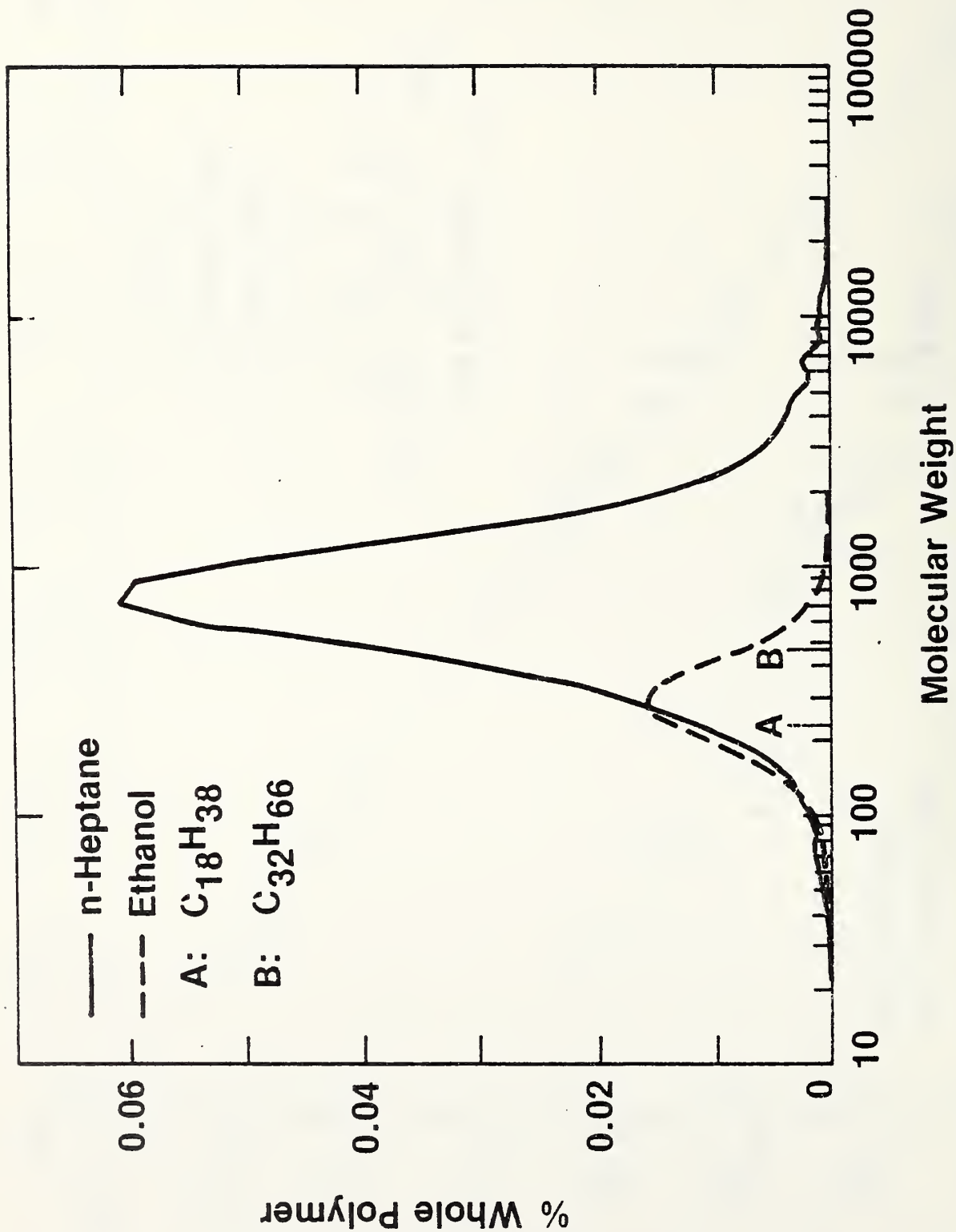


Figure VIII-4 Molecular Weight Distributions of Ethanol and n-Heptane Extracts of LPE

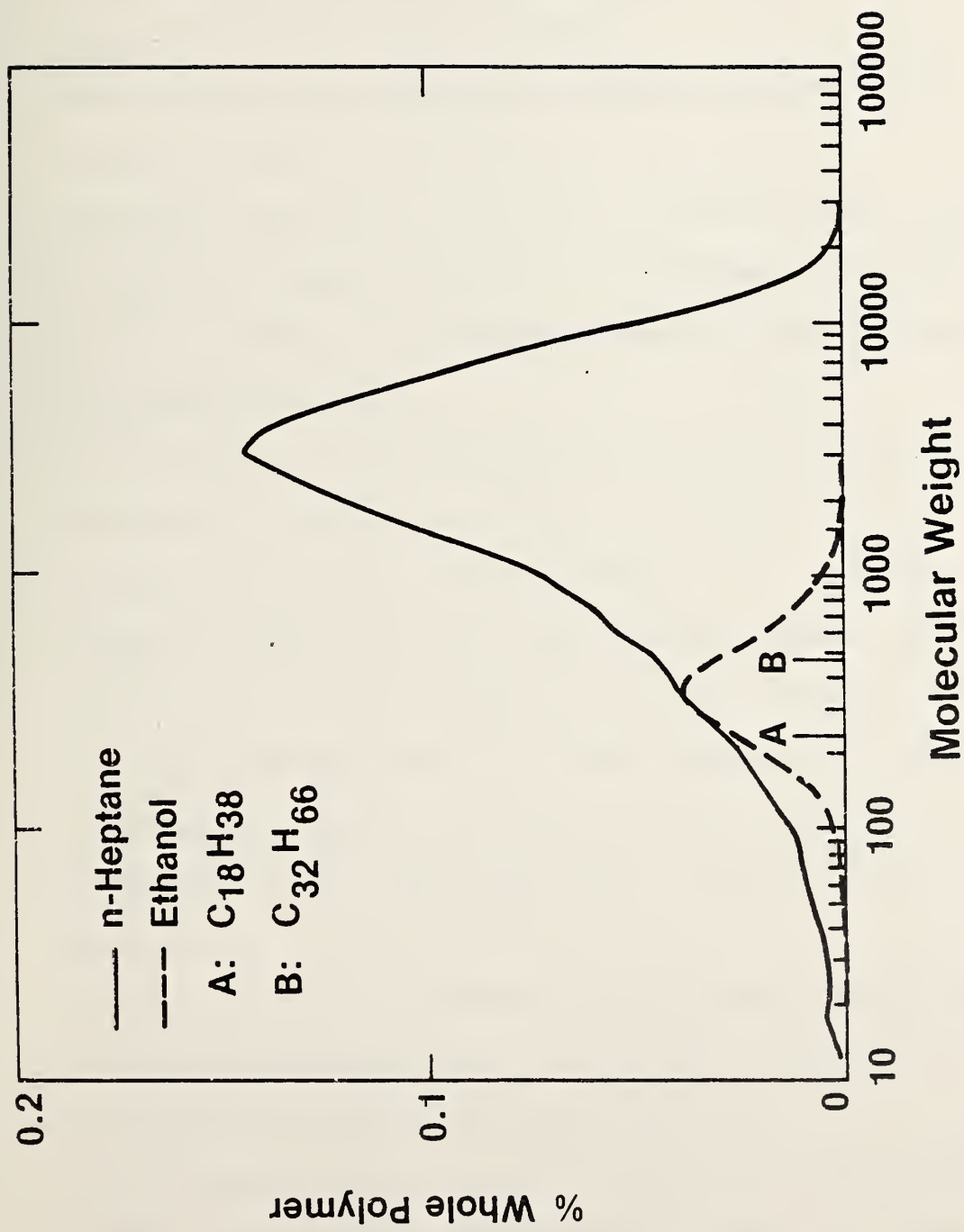
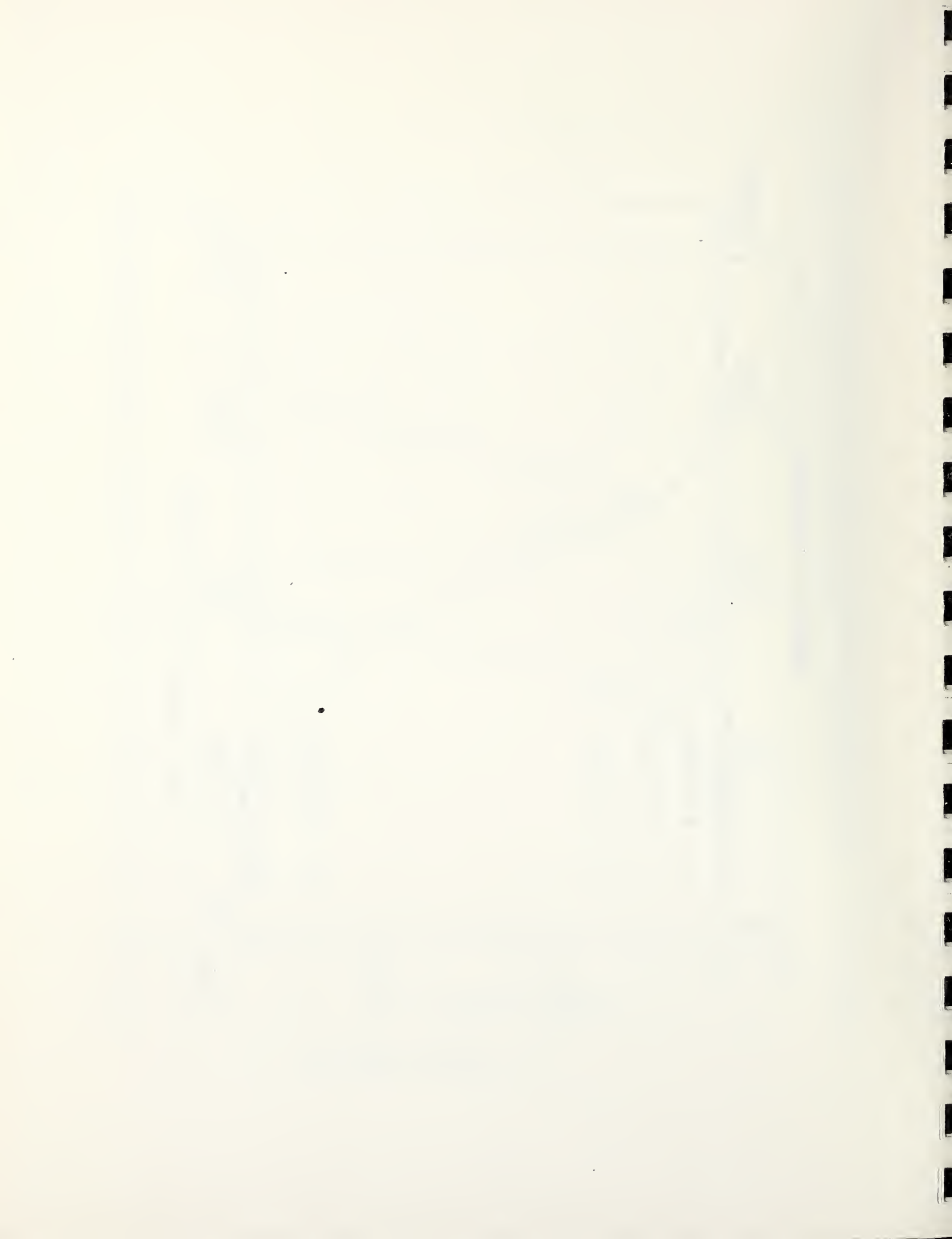


Figure VIII-5 Molecular Weight Distributions of Ethanol and n-Heptane Extracts of BPE



IX. Summary of Conclusions

- * Maximum amount of extractables depends on the original migrant concentration in the polymer, the solubilities and the partition of the migrant in solvent and in polymer.
- * Migration behavior is mainly Fickian.
- * Diffusion coefficient may be increased by absorption of the solvent or swelling of the polymer by the solvent.
- * Activation energies for the change of diffusion coefficient as a function of temperature generally lie between 40 and 200 kJ mol⁻¹. In general, the activation energy is lower at higher diffusion coefficients.
- * Lower members of pure triglycerides may be used to simulate the extraction behavior of corn oil in oligomers and anti-oxidants, such as BHT, from polyolefins.
- * Anhydrous ethanol may also be considered as food-oil simulant.
- * Accelerating action of n-heptane over that of food-oils or oil simulants is more pronounced at lower diffusion coefficients.
- * n-Heptane will remove high molecular weight fractions of oligomers which may hardly be present in food oil or oil simulants.
- * Diffusion coefficient increases with increased oligomer content.
- * Increase of amorphous content in a polyolefin produces a corresponding change in diffusion coefficient.
- * The following is the ranking of the importance of the parameters effecting the diffusion coefficient of a migrant from polymer matrix into surrounding media:

Solvent > Temperature (limited range) > Swelling

> Migrant Concentration > Amorphous content.



NBSIR 82-2472

Migration of Low Molecular Weight Additives
in Polyolefins and Copolymers

X. Appendix

Numerical Data of Migration of Low Molecular
Weight Additives from Polyolefins and Copolymers

Experimental Run Designations--cf page II-6, Tables II-4 and II-5



LPE/C18/CO/30C/L	t, h	Mt/Mo	
0.033	0.0055	0.0055	
0.167	0.00296	0.00296	
0.333	0.00396	0.00396	
0.5	0.00462	0.00462	
1.17	0.0115	0.0115	
2.4	0.0202	0.0202	
4.5	0.028	0.028	
7.8	0.0422	0.0422	
12.5	0.0494	0.0494	
21.3	0.0556	0.0556	
30.8	0.103	0.103	
45.6	0.142	0.142	
51.1	0.175	0.175	
67.5	0.245	0.245	
96.6	0.314	0.314	
125	0.372	0.372	
158.5	0.412	0.412	
216.4	0.505	0.505	
285.7	0.572	0.572	
362.5	0.634	0.634	
413.2	0.681	0.681	
482.5	0.71	0.71	
552.3	0.739	0.739	
626.8	0.765	0.765	
694.0	0.789	0.789	
765.7	0.804	0.804	
821.3	0.815	0.815	
873.7	0.824	0.824	
	0.822	0.822	

LPE/C18/CO/60C/L

	t, h	Mt/Mo	
	0.017	0.0104	
	0.033	0.019	
	0.067	0.0269	
	0.133	0.0378	
	0.25	0.056	
	0.5	0.0855	
	1.5	0.125	
	2.5	0.15	
	4	0.227	
	6	0.268	
	8	0.304	
	23.5	0.47	
	31	0.524	
	48.5	0.631	
	54	0.651	
	120	0.834	
	128	0.85	
	198	0.914	
	289	0.959	
	457	0.985	
	725	0.996	
	965	0.981	

LPE/C18/EH/60C/L
120600

t, h	Mt/Mo
0.017	0.0149
0.033	0.0185
0.067	0.0287
0.133	0.0456
0.25	0.0693
0.5	0.0996
1.5	0.135
2.5	0.159
4.5	0.197
7.5	0.239
12.5	0.275
24.5	0.323
55.8	0.527
120	0.745
216	0.92
384	0.98
384	1.06
558	0.997
558	0.961
	0.957

LPE/C18/.5EN/60C/L(1)

LPE/C18/.5EN/60C/L(2)

LPE/C18/.5EN/60C/L(3)

t, h	Mt/Mo	t, h	Mt/Mo
0.017	9.16E-4	0.017	6.69E-4
0.033	0.00176	0.033	0.00126
0.067	0.0028	0.067	0.00241
0.133	0.00485	0.133	0.00438
0.25	0.00821	0.25	0.00741
0.5	0.0147	0.5	0.015
1	0.0263	1	0.0263
1.5	0.0342	1.5	0.0434
2.5	0.0471	2.5	0.0609
4	0.0571	4	0.0964
5.5	0.0635	5.5	0.0962
7	0.0677	7	0.107
23.8	0.0683	23.8	0.117

t, h	Mt/Mo
0.017	6.69E-4
0.033	0.00126
0.067	0.00241
0.133	0.00438
0.25	0.00741
0.5	0.015
1	0.0263
1.5	0.0434
2.5	0.0609
4	0.0964
5.5	0.107
7	0.117

127600
LPE/C18/.7EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00912	0.017	0.0172
0.033	0.0161	0.033	0.0215
0.067	0.0251	0.067	0.0299
0.133	0.0341	0.133	0.0443
0.25	0.0505	0.25	0.063
0.5	0.0771	0.5	0.0903
1.5	0.133	1.5	0.126
2.5	0.163	2.5	0.148
4.5	0.196	4.5	0.182
7.5	0.22	7.5	0.221
24.5	0.383	24.5	0.25
30	0.429	30	0.456
96.5	0.565	96.5	0.515
147	0.568	147	0.616
196	0.578	196	0.658
264	0.569	264	0.818
344	0.577	344	0.829
			0.87
			0.885
			0.886
			0.893
			0.89
			0.905

129600
LPE/C18/.9EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0172	0.017	0.0172
0.033	0.0215	0.033	0.0215
0.067	0.0299	0.067	0.0299
0.133	0.0443	0.133	0.0443
0.25	0.063	0.25	0.063
0.5	0.0903	0.5	0.0903
1.5	0.126	1.5	0.126
2.5	0.148	2.5	0.148
4.5	0.182	4.5	0.182
7.5	0.221	7.5	0.221
24.5	0.25	24.5	0.25
30	0.456	30	0.456
96.5	0.515	96.5	0.515
147	0.616	147	0.616
196	0.658	196	0.658
264	0.818	264	0.818
344	0.829	344	0.829
	0.87		0.87
	0.885		0.885
	0.886		0.886
	0.893		0.893
	0.89		0.89
	0.905		0.905

130240
LPE/C18/HP/24C/L130301
LPE/C18/HP/30C/U130601
LPE/C18/HP/60C/U1

t, h	Mt/Mo
0.1	0.0456
16	0.836
23	0.891
40	0.932
70	0.955
135	0.964
216	0.969
309	0.972
358	0.975
666	0.974

130241
LPE/C18/HP/24C/U

t, h	Mt/Mo
0.025	0.0262
0.05	0.0392
0.1	0.0533
0.25	0.0827
0.5	0.115
1	0.163
2	0.242
4	0.378
8	0.662
12	0.902
24	0.947
72	0.952
151	0.958
245	0.971
294	0.974
438	0.971
462	0.975
512	0.975
579	0.975

t, h	Mt/Mo
0.1	0.017
16	0.033
23	0.067
40	0.133
70	0.267
135	0.5
216	0.75
309	1.5
358	2.5
666	4

t, h	Mt/Mo
0.1	0.0226
16	0.0348
23	0.0463
40	0.0652
70	0.0914
135	0.123
216	0.151
309	0.176
358	0.219
666	0.295

t, h	Mt/Mo
0.1	0.25
16	0.5
23	1.2
40	2.4
70	4.8
135	12.0
216	16.8

t, h	Mt/Mo
0.1	0.25
16	0.5
23	1.2
40	2.4
70	4.8
135	12.0
216	16.8

140301 LPE/C18/0D/30C/U
 t, h Mt/Mo
 0.017 0.00733
 0.033 0.0114
 0.067 0.0163
 0.133 0.0263
 0.267 0.0418
 0.5 0.0593
 0.75 0.0733
 1.5 0.0844
 2.5 0.102
 4.5 0.126
 7.5 0.157
 12.5 0.182
 24.5 0.21
 31.5 0.24
 96 0.376
 103 0.437
 103 0.865
 145 0.887
 145 0.936
 199 0.949
 264 0.955
 320 0.958
 439 0.962
 534 0.964
 672 0.967
 1033 0.971
 1372 0.974
 1561 0.974

140601 LPE/C18/0D/60C/U
 t, h Mt/Mo
 0.017 0.017
 0.033 0.033
 0.067 0.067
 0.133 0.133
 0.25 0.25
 0.5 0.5
 1.2 1.2
 4 4
 6.08 6.08
 7.5 7.5
 24 24

LPE/C18/T0/300C/L

LPE/C18/T0/600C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 017	0. 00196	0. 017	0. 0137
0. 033	0. 00243	0. 033	0. 0217
0. 067	0. 0038	0. 067	0. 0317
0. 133	0. 00597	0. 133	0. 0462
0. 25	0. 00878	0. 25	0. 0676
0. 5	0. 013	0. 5	0. 0969
1. 5	0. 0208	1. 5	0. 131
2. 5	0. 027	2. 5	0. 155
4. 5	0. 0365	4. 5	0. 192
7. 5	0. 0482	7. 5	0. 228
14. 5	0. 056	14. 5	0. 266
24. 5	0. 0648	24. 5	0. 288
49. 5	0. 0713	49. 5	0. 345
98. 5	0. 08143	98. 5	0. 422
197. 5	0. 0925	197. 5	0. 47
385	0. 09324	385	0. 583
703	0. 09345	703	0. 651
894	0. 0947	894	0. 715
1513	0. 0959	1513	0. 768
2092	0. 0961	2092	0. 859
2785	0. 0963	2785	0. 863
3553	0. 0964	3553	0. 864
4061	0. 0965	4061	0. 861
4753	0. 0966	4753	0. 882
5451	0. 0967	5451	0. 841
6196	0. 0968	6196	0. 859
6868	0. 0969	6868	0. 863
7585	0. 0970	7585	0. 864
8146	0. 0971	8146	0. 882
8740	0. 0972	8740	0. 882

220240
LPE/C18/EN/24C/L

	t, h	Mt/Mo	t, h	Mt/Mo
1	0.114	0.017	0.00744	
3	0.161	0.033	0.0127	
5	0.335	0.067	0.0173	
116	0.639	0.133	0.0252	
151.5	0.735	0.25	0.0384	
195	0.778	0.5	0.0577	
288	0.799	1.5	0.0807	
337	0.816	2.5	0.114	
387	0.832	4.5	0.149	
505	0.857	7	0.186	
643	0.855	12	0.217	
820	0.883	23	0.241	
1220	0.9	31	0.29	
1556		48	0.441	

220300
LPE/C18/EN/30C/L

	t, h	Mt/Mo	t, h	Mt/Mo
1	0.114	0.017	0.00744	
3	0.161	0.033	0.0127	
5	0.335	0.067	0.0173	
116	0.639	0.133	0.0252	
151.5	0.735	0.25	0.0384	
195	0.778	0.5	0.0577	
288	0.799	1.5	0.0807	
337	0.816	2.5	0.114	
387	0.832	4.5	0.149	
505	0.857	7	0.186	
643	0.855	12	0.217	
820	0.883	23	0.241	
1220	0.9	31	0.29	
1556		48	0.441	

220241
LPE/C18/EN/24C/U

	t, h	Mt/Mo	t, h	Mt/Mo
0.083	0.0165	0.0165	0.0165	
0.167	0.0234	0.0234	0.0234	
0.33	0.0352	0.0352	0.0352	
0.67	0.0518	0.0518	0.0518	
1	0.0686	0.0686	0.0686	
2	0.109	0.109	0.109	
4	0.166	0.166	0.166	
7	0.237	0.237	0.237	
24	0.367	0.367	0.367	
48	0.465	0.465	0.465	
78	0.578	0.578	0.578	
216	0.729	0.729	0.729	
409	0.809	0.809	0.809	
745	0.865	0.865	0.865	

220600
LPE/C18/EN/60C/L220601
LPE/C18/EN/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.05	0.017	0.0708
0.033	0.0708	0.05	0.135
0.067	0.101	0.133	0.232
0.133	0.157	0.3	0.331
0.25	0.239	0.55	0.423
0.5	0.35	1.05	0.543
1.5	0.572	2.2	0.688
2.5	0.698	4.5	0.84
4.5	0.808	7.5	0.924
5.52	0.854	24	0.971
7.92	0.911	25	0.98
23.6	0.968	147	0.981
31	0.966	319	0.983
55.8	0.968	414	0.986
122	0.965	532	0.987
220	0.929	721	0.988
384	0.971	721	0.979
626	0.968		
721	0.969		
721	0.97		

225600
LPE/C18/.5EN/60C/L(1)

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00314	1.15	0.105
0.033	0.0053	2.65	0.146
0.067	0.00913	5.98	0.157
0.133	0.0169	71.3	0.165
0.25	0.0327	168	0.156
0.5	0.0603	240	0.111
0.75	0.0814		
1.1	0.0971		
1.5	0.119		
2.5	0.152		
4.5	0.164		
7.5	0.173		
23.7	0.172		
31.5	0.172		
54.5	0.174		
168	0.176		
344	0.178		

225603
LPE/C18/.5EN/60C/L(3)

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00314	1.15	0.105
0.033	0.0053	2.65	0.146
0.067	0.00913	5.98	0.157
0.133	0.0169	71.3	0.165
0.25	0.0327	168	0.156
0.5	0.0603	240	0.111
0.75	0.0814		
1.1	0.0971		
1.5	0.119		
2.5	0.164		
4.5	0.173		
7.5	0.172		
23.7	0.172		
31.5	0.172		
54.5	0.174		
168	0.176		
344	0.178		

LPE/C18/.5EN/60C/L(4)

t, h	Mt/Mo	t, h	Mt/Mo
0.967	0.0639		
3.22	0.0906		
6.13	0.0954		
24	0.0981		
47.9	0.0971		
268	0.0987		
409	0.0978		

LPE/C18/.5EN/60C/L(5)

t, h	Mt/Mo	t, h	Mt/Mo
0.167	0.0107		
0.333	0.0198		
0.667	0.0366		
1	0.0510		
2	0.0664		
4	0.08		
5.95	0.0744		
22.9	0.0758		
29.9	0.0773		
94.9	0.0714		
195	0.0314		

LPE/C18/.5EN/60C/U
225601

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00936	1.5	0.191
0.033	0.0161	2.5	0.255
0.067	0.0267	3.5	0.327
0.133	0.0488	4.5	0.409
0.25	0.0806	7.5	0.488
0.5	0.119	24.5	0.997
1.1		30.5	0.652
2.2		120	0.823
4.4		361	0.926
7.7		698	0.964
24		1012	0.965
31		1204	0.978
48		1347	0.981
71			
96			
120			
193			
222			
295			
390			
508			
519			

LPE/C18/.9EN/60C/U
229601

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	1	0.051
0.033	0.033	2	0.058
0.067	0.083	4	0.087
0.133	0.167	7	0.093
0.25	0.25	24	0.094
0.5	0.5	31	0.097
1.1		48	0.096
2.2		71	0.098
4.4		96	0.098
7.7		120	0.098
24		193	0.0983
31		222	0.0984
48		295	0.0984
71		390	0.0985
96		508	0.0986
120		519	0.0988

LPE/C18/HP/24C/U

LPE/C18/HP/30C/U

t, h

Mt/Mo

0.017 0.0857

0.033 0.124

0.067 0.183

0.133 0.281

0.25 0.458

0.5 0.683

1 0.804

2 0.866

4 0.904

6 0.919

8 0.955

10 0.959

12 0.972

14 0.974

16 0.975

18 0.994

20 0.994

22 0.994

24 0.997

26 0.997

28 0.997

30 0.997

32 0.997

34 0.997

36 0.997

38 0.997

LPE/C18/00/60C/U

LPE/C18/00/24C/U

t, h

Mt/Mo

0.017 0.017

0.033 0.033

0.067 0.067

0.133 0.133

0.25 0.25

0.5 0.5

1 1

2 2

4 4

6 6

8 8

10 10

12 12

14 14

16 16

18 18

20 20

22 22

24 24

26 26

28 28

30 30

32 32

34 34

LPE/C18/00/24C/U

LPE/C18/00/60C/U

t, h

Mt/Mo

0.017 0.0736

0.033 0.119

0.067 0.19

0.133 0.3

0.25 0.472

0.5 0.715

1 0.812

2 0.844

4 0.874

6 0.906

8 0.917

10 0.939

12 0.959

14 0.962

16 0.966

18 0.968

20 0.969

22 0.971

24 0.974

26 0.977

28 0.979

30 0.981

32 0.984

34 0.987

LPE/C18/00/60C/U

LPE/C18/00/24C/U

t, h

Mt/Mo

0.017 0.25

0.033 0.5

0.067 1.2

0.133 2.4

0.25 4.8

0.5 127

1 168

LPE/C18/0H/30C/L

250300
LPE/C18/0H/60C/L

t, h	Ht/Mo						
0.017	0.0132	0.017	0.0132	0.017	0.0132	0.017	0.0132
0.033	0.011	0.057	0.0128	0.033	0.011	0.057	0.0128
0.133	0.018	0.267	0.0294	0.133	0.018	0.267	0.0294
0.267	0.05	0.464	0.0464	0.267	0.05	0.464	0.0464
2.03	0.141	3.5	0.0965	2.03	0.141	3.5	0.0965
5.7	0.189	6.5	0.0278	5.7	0.189	6.5	0.0278
24.7	0.49	24.7	0.233	24.7	0.49	24.7	0.233
39.5	0.541	48.8	0.652	39.5	0.541	48.8	0.652
76.8	0.77	145	0.834	76.8	0.77	145	0.834
145	0.87	241	0.874	145	0.87	241	0.874
241	0.874	337	0.874	241	0.87	337	0.874
484	0.899	649	0.913	484	0.899	649	0.913
676	0.916			676	0.916		

LPE/310300

LPE/C18/C0/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.25	7.95E-4		
0.5	0.00127		
1	0.00258		
1.5	0.00376		
2.5	0.00558		
4	0.0066		
6.5	0.00793		
9	0.01199		
12	0.0241		
15	0.0285		
24	0.04529		
32	0.0609		
49	0.0763		
59	0.0877		
75	0.107		
84	0.116		
92	0.116		
120	0.133		
156	0.15		
192	0.166		
239	0.186		
290	0.205		
341	0.22		
391	0.234		
437	0.249		
487	0.268		
545	0.278		
603	0.295		
653	0.308		
705	0.319		
787	0.336		
842	0.346		
877	0.353		

t, h	Mt/Mo	t, h	Mt/Mo
0.25	0.017	0.00268	
0.5	0.033	0.00447	
1	0.067	0.00707	
1.5	0.133	0.013	
2.5	0.25	0.0208	
4	0.5	0.0387	
6.5	1.5	0.0743	
9	2.5	0.0936	
12	5	0.117	
15	5.5	0.1555	
24	8.5	0.2295	
32	12.5	0.385	
55	24.5	0.46	
81	37.5	0.592	
150	73	0.73	
240	150	0.815	
337	240	0.883	
488	337	0.935	
721	488	0.971	
1154	721	0.996	
1400	1154	0.999	
1584	1400	0.999	
1920	1584	0.999	
2404	1920	0.983	
3030	2404	0.942	
3838	3030	0.944	

320600
LPE/C18/EN/60C/L

t, h	Mt/Mo
0.017	0.00284
0.033	0.00434
0.067	0.00758
0.133	0.0155
0.25	0.0275
0.5	0.0439
1.5	0.0833
2.5	0.162
4	0.129
5.5	0.149
8	0.175
12.5	0.206
24	0.329
32	0.437
56	0.58
105	0.701
170	0.762
241	0.798
344	0.831
505	0.852
720	0.873
1086	0.879
1273	0.872
1583	0.878
1945	0.886
1947	0.892
2400	0.891
2400	0.895
2883	0.899
2883	0.805

LPE/C18/.1EH/60C/L

321600
LPE/C18/.5EH/60C/L(1)

t, h	Mt/Mo
24	8.42E-5
96	2.2E-4
192	2.63E-4
528	2.86E-4
792	2.72E-4
1464	2.79E-4

323600
LPE/C18/.3EH/60C/L

t, h	Mt/Mo
24	4.09E-4
96	7.31E-4
192	7.28E-4
528	7.03E-4
792	7.05E-4
1464	6.57E-4

325602
LPE/C18/.5EH/60C/L(2)

t, h	Mt/Mo
24	0.017
96	0.067
192	0.167
528	0.333
792	1.07
1464	2
2304	4
3300	7
4300	23.7
5300	30
6300	95.8
7300	193
8300	340
9300	533
10300	672

330241 LPE/C18/HP/24C/U

330301 LPE/C18/HP/30C/U

330601 LPE/C18/HP/68C/U

t, h Mt/Mo

0.067 0.0153
0.133 0.0203
0.25 0.0286
0.5 0.0393
1.2 0.0559
2.4 0.0856
8 0.128
71.3

244
293
317
461
511
604
775
1008

t, h Mt/Mo

0.033 0.00727
0.067 0.0108
0.133 0.0169
0.25 0.0248
0.5 0.0382
1.2 0.0584
2.4 0.0757
5 0.0815
5.5 0.0856
11.5 0.128
19.6 0.196
46.7 0.286
66.3 0.394
84.8 0.48
85.3 0.53
48 0.56
56 0.69
67.1 0.78
69.4 0.87
78 0.94
87.3 0.97
88.7 0.98
89.4 0.

126
175
175
1224
224
337
489
720
871
11536
1204
1901
2407
2905
3409
3894
4517

t, h Mt/Mo

0.033 0.00727
0.067 0.0108
0.133 0.0169
0.25 0.0248
0.5 0.0382
1.2 0.0584
2.4 0.0757
5 0.0815
5.5 0.0856
11.5 0.128
19.6 0.196
46.7 0.286
66.3 0.394
84.8 0.48
85.3 0.53
48 0.56
56 0.69
67.1 0.78
69.4 0.87
78 0.94
87.3 0.97
88.7 0.98
89.4 0.

t, h Mt/Mo

0.033 0.00727
0.067 0.0108
0.133 0.0169
0.25 0.0248
0.5 0.0382
1.2 0.0584
2.4 0.0757
5 0.0815
5.5 0.0856
11.5 0.128
19.6 0.196
46.7 0.286
66.3 0.394
84.8 0.48
85.3 0.53
48 0.56
56 0.69
67.1 0.78
69.4 0.87
78 0.94
87.3 0.97
88.7 0.98
89.4 0.

t, h Mt/Mo

0.033 0.00727
0.067 0.0108
0.133 0.0169
0.25 0.0248
0.5 0.0382
1.2 0.0584
2.4 0.0757
5 0.0815
5.5 0.0856
11.5 0.128
19.6 0.196
46.7 0.286
66.3 0.394
84.8 0.48
85.3 0.53
48 0.56
56 0.69
67.1 0.78
69.4 0.87
78 0.94
87.3 0.97
88.7 0.98
89.4 0.

LPE/C18/00/30C/U

0.083	0.00473
0.167	0.0087
0.333	0.0145
0.667	0.0228
1.2	0.0303
2.4	0.0484
4.7	0.0772
7.3	0.112
10.3	0.241
14.4	0.295
19.9	0.355
26.7	0.415
36.0	0.475
45.6	0.534
69.6	0.601
127.2	0.661
161.3	0.720
194.2	0.780
238.0	0.840
288.6	0.905
350.7	0.91

LPE/C18/00/60C/U

t, h	Mt/Mo
0.033	0.0411
0.067	0.0628
0.133	0.0945
0.25	0.136
0.5	0.196
1.1	0.285
1.5	0.352
2.5	0.463
4.5	0.595
8.0	0.708
12.4	0.889
23.2	0.923
57	0.933
122	0.943
176	0.954
225	0.958
344	0.964
481	0.967
720	0.97
1038	0.972
1362	0.974
1560	0.975
1900	0.977
2405	0.979

370300
LPE/C18/T0/30C/LLPE/C18/T0/60C/L
370600

t, h	Mt/Mo	t, h	Mt/Mo
0.017	8.91E-4	0.017	0.00409
0.033	8.93E-4	0.033	0.00611
0.067	8.71E-4	0.067	0.00857
0.133	0.00103	0.133	0.0153
0.25	0.00144	0.25	0.0268
0.5	0.00184	0.5	0.0441
1.1	0.00256	1.1	0.067
2.5	0.00362	2.5	0.0817
5	0.00517	5	0.104
11.5	0.00763	11.5	0.131
25	0.0114	25	0.153
48	0.0241	48	0.181
120	0.0333	120	0.296
336	0.0359	336	0.339
480	0.0509	480	0.563
702	0.0597	702	0.72
872	0.0816	872	0.837
1192	0.0988	1192	0.876
1540	0.116	1540	0.918
1969	0.129	1969	0.94
2399	0.147	2399	0.949
2908	0.169	2908	0.944
3408	0.189	3408	0.942
3888	0.208	3888	0.942
4343	0.223	4343	0.942
4830	0.245	4830	0.942
5411	0.262	5411	0.942
5982	0.279	5982	0.942
6482	0.302	6482	0.942
7914	0.309	7914	0.942
8425	0.328	8425	0.942
8957	0.341	8957	0.942
9425	0.354	9425	0.942
9957	0.376	9957	0.942
0.388	0.393	0.388	0.942

380600
LPE/C18/AQ/600/L

t, h	n _t /n ₀
149	0.00147
216	0.00261
389	0.00707
624	0.0165
866	0.0268
1102	0.0352
11538	0.0522
1968	0.0642
2357	0.0743
2909	0.0888
3480	0.0986
4060	0.106
4705	0.117
5335	0.12

430241
LPE/C18/HP/24C/U

t, h	Mt/Mo
0.033	0.0148
0.067	0.0258
0.133	0.0411
0.25	0.0628
0.5	0.108
1	0.202
2	0.342
4	0.477
8	0.693
16	0.732
32	0.817
64	0.838
128	0.851
256	0.861
512	0.873
1024	0.865

LPE/C18/HP/30C/U
530301

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0834	0.167	0.129
0.033	0.143	0.333	0.185
0.083	0.184	0.667	0.264
0.167	0.279	1	0.319
0.333	0.385	2	0.426
0.5	0.458	4	0.561
0.75	0.542	8	0.714
1	0.612	24	0.94
1.25	0.678	48	0.991
1.75	0.772	79.5	0.997
2.25	0.925	169	0.999
2.75	0.947	217	0.999
3.75	0.96	389	1
4.75	0.979		
5.75	0.981		
7.75	0.987		
11.75	0.988		
14.75	0.989		
19.9	0.989		
26.5	0.99		
43.9	0.991		
53.4	0.991		
65.2	0.991		
66.3	0.994		

LPE/C18/OD/30C/U
540301

t, h	Mt/Mo	t, h	Mt/Mo
0.167	0.169	0.167	0.169
0.333	0.278	0.333	0.278
0.667	0.404	0.667	0.404
1	0.493	1	0.493
1.5	0.601	1.5	0.601
2.5	0.734	2.5	0.734
4	0.864	4	0.864
6	0.937	6	0.937
8	0.971	8	0.971
23	0.994	23	0.994
231.5	0.995	231.5	0.995
118.9	0.995	118.9	0.995
227	0.996	227	0.996
311	0.996	311	0.996

640301
LPE/C18/00/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.012	0.109	0.017	0.128
0.025	0.155	0.033	0.18
0.05	0.219	0.067	0.254
0.09	0.317	0.083	0.285
0.104	0.361	0.133	0.405
0.133	0.405	0.167	0.524
0.167	0.524	0.25	0.599
0.25	0.599	0.337	0.703
0.337	0.703	0.594	0.784
0.594	0.784	0.667	0.875
0.667	0.875	0.1	0.934
0.1	0.934	1.5	0.955
1.5	0.955	2.3	0.968
2.3	0.968	4.5	0.972
4.5	0.972	6	0.975
6	0.975	8	0.976
8	0.976	25.3	0.981

640601
LPE/C18/00/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.012	0.376	0.017	0.582
0.025	0.772	0.033	0.923
0.05	0.921	0.067	0.981
0.09	0.992	0.083	0.992
0.104	0.992	0.167	0.992
0.133	0.992	0.333	0.992
0.167	0.992	0.667	0.992
0.25	0.992	1.5	0.992
0.337	0.992	2	0.992
0.594	0.992	3	0.992
0.667	0.992	4	0.992
0.1	0.992	5	0.992
1.5	0.992	7	0.992
2	0.992	7	0.992
3	0.992	7	0.992
4	0.992	7	0.992
5	0.992	7	0.992
6	0.992	7	0.992
8	0.992	7	0.992
19.9	0.997	19.9	0.997

710301
BPE/C18/CO/38C/u

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.0115	0.017	0.0332
0.067	0.0215	0.033	0.0641
0.133	0.0323	0.067	0.093
0.25	0.044	0.133	0.134
0.5	0.0581	0.25	0.188
1.5	0.0749	1.5	0.267
2.5	0.0895	3.5	0.368
4.5	0.109	5.5	0.447
8.5	0.131	11.5	0.745
24	0.15	24	0.919
48	0.174	48	0.945
81	0.258	81	0.958
144	0.341	144	0.97
241	0.425	241	0.98
389	0.532	389	0.99
682	0.624	682	0.999
1181	0.708	1181	0.999
1543	0.769	1543	0.999
865	0.817	865	0.999
1921	0.849	1921	0.999
2466	0.873	2466	0.999
2885	0.908	2885	0.999
3779	0.922	3779	0.999
4284	0.938	4284	0.999
4854	0.95	4854	0.999
5407	0.96	5407	0.999

710601
BPE/C18/CO/60C/u

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0115	0.017	0.0332
0.033	0.0215	0.033	0.0641
0.067	0.0323	0.067	0.093
0.133	0.044	0.133	0.134
0.25	0.0581	0.25	0.188
0.5	0.0749	0.5	0.267
1.5	0.0895	1.5	0.368
3.5	0.109	5.5	0.447
11.5	0.131	24	0.745
24	0.15	24	0.919
48	0.174	48	0.945
81	0.258	81	0.958
144	0.341	144	0.97
241	0.425	241	0.98
389	0.532	389	0.99
682	0.624	682	0.999
1181	0.708	1181	0.999
1543	0.769	1543	0.999
865	0.817	865	0.999
1921	0.849	1921	0.999
2466	0.873	2466	0.999
2885	0.908	2885	0.999
3779	0.922	3779	0.999
4284	0.938	4284	0.999
4854	0.95	4854	0.999
5407	0.96	5407	0.999

720301
BPE/C18/EN/30C/U720601
BPE/C18/EN/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.00813	0.033	0.017
0.067	0.0106	0.033	0.0527
0.133	0.0145	0.067	0.0743
0.25	0.0199	0.133	0.104
0.5	0.0286	0.25	0.146
1.	0.0419	0.5	0.199
1.5	0.0519	1.	0.286
2.5	0.0694	2.5	0.582
4.5	0.0903	4.5	0.779
5.	0.100	5.	0.877
24.3	0.172	24.	0.917
48.	0.244	3	0.999
120	0.356	98.	0.998
216	0.556	509	0.999
385	0.689	1277	0.999
555	0.789		
866	0.838		
1181	0.881		
1537	0.904		
2022	0.919		
2401	0.932		
2881	0.939		
3339	0.945		
	0.95		

730301
BPE/C18/HP/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.0457	0.033	0.16
0.067	0.0692	0.033	0.229
0.133	0.106	0.067	0.336
0.25	0.157	0.133	0.484
0.5	0.243	0.25	0.693
1	0.376	0.5	0.932
1.5	0.499	1	0.996
2.5	0.72	1.5	0.999
4	0.862	2.5	0.999
5.5	0.912	4.5	1
8	0.942	5.5	1
24	0.974	8	1
32	0.977	24	1
48	0.98	32	1
119	0.986	48	1
215	0.988	119	1
383	0.989	215	1
628	0.99	383	1

730601
BPE/C18/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.017	0.033	0.16
0.067	0.033	0.067	0.229
0.133	0.067	0.133	0.336
0.25	0.133	0.25	0.484
0.5	0.25	0.5	0.693
1	0.5	0.5	0.932
1.5	1	0.996	1
2.5	1.5	0.999	1
4	2.5	0.999	1
5.5	4.5	0.999	1
8	5.5	0.999	1
24	8	0.999	1
32	24	0.999	1
48	32	0.999	1
119	48	0.999	1
215	119	0.999	1
383	215	0.999	1
628	383	0.999	1

770301
BPE/C18/T0/300/U

770601
BPE/C18/T0/600/U

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.0127	0.017	0.0448
0.067	0.0224	0.033	0.0731
0.133	0.0323	0.067	0.109
0.25	0.0434	0.133	0.154
0.5	0.0571	0.25	0.208
1.5	0.0738	0.5	0.286
2.5	0.0993	1.02	0.397
4.5	0.108	1.5	0.475
9.5	0.13	2.5	0.592
24.5	0.154	4	0.716
48.2	0.264	5.5	0.793
76.5	0.353	24	0.856
96.	0.429	31.5	0.934
216	0.474	71.3	0.946
386	0.626	169	0.96
605	0.724	748	0.972
870	0.785	1537	0.984
1177	0.824	11	0.997
1536	0.87		
1950	0.895		
2406	0.898		
2888	0.908		
3368	0.917		
3701	0.932		

BPE/C18/HIP/30C/U

t, h	Mt/Mo
0.006	0.0452
0.008	0.057
0.017	0.0859
0.033	0.129
0.067	0.187
0.133	0.274
0.25	0.388
0.5	0.582
1	0.814
1.5	0.901
2.5	0.955
4	0.976
5.67	0.982
7	0.984
23.8	0.99
31.3	0.991
48	0.992
74.9	0.993
173	0.994

840301
BPE/C18/0D/30C/U1

840302
BPE/C18/0D/30C/U2

840601
BPE/C18/0D/60C/U

t, h	Mt/Mo	BPE/C18/0D/30C/U1	BPE/C18/0D/30C/U2	BPE/C18/0D/60C/U
0.004	0.0225	0.011	0.037	0.0654
0.008	0.0402	0.019	0.055	0.104
0.021	0.0587	0.025	0.077	0.143
0.033	0.0737	0.033	0.098	0.166
0.05	0.0905	0.053	0.119	0.198
0.083	0.117	0.067	0.133	0.218
0.117	0.14	0.077	0.147	0.248
0.183	0.176	0.089	0.126	0.305
0.233	0.197	0.099	0.147	0.391
0.35	0.24	0.103	0.172	0.476
0.5	0.2795	0.117	0.225	0.544
0.983	0.34	0.133	0.253	0.652
1.17	0.484	0.183	0.333	0.77
1.4	0.55	0.253	0.5	0.848
1.98	0.649	0.333	0.75	0.932
2.93	0.721	0.415	1.5	0.969
3.98	0.82	0.477	2.2	0.992
5.98	0.904	0.577	3.5	0.998
8.98	0.987	0.694	5.5	0.999
23.8	0.991	0.771	8	0.999
32.8	0.993	0.852	24	0.999
48.2	0.993	0.981	32	0.999
56.4	0.993	0.987	48	0.999
72.3	0.994	0.989	56	0.999
80.3	0.994	0.999	73.5	0.999
169	0.995	0.995	73.5	0.999
173				0.999

920300
BPE/C18/EN/30C/L
t, h Mt/Mo

920600
BPE/C18/EN/60C/L
t, h Mt/Mo

0.067	0.0086	0.017	0.053
0.133	0.0166	0.033	0.0781
0.25	0.0196	0.067	0.0955
0.5	0.0317	0.133	0.127
1.97	0.0479	0.25	0.177
4.25	0.094	0.5	0.273
24.	0.231	1.5	0.404
30.8	0.271	2.5	0.641
55.8	0.391	4	0.802
96.3	0.511	5	0.884
144	0.555	6	0.959
193	0.633	7	1.04
264	0.713	8	1.04
316	0.728	9	1.04
361	0.767	10	1.04
433	0.765	11	1.04
481	0.791	12	1.04
601	0.823	13	1.04
701		14	1.04

925380
BPE/C18/.5EN/30C/L

925600
BPE/C18/.5EN/60C/L

t, h Mt/Mo t, h Mt/Mo

2.5 0.00399
4 0.00655
5.78 0.0102
8 0.0101
23.6 0.0151
31.5 0.0158
348.1 0.0163
55.7 0.0158
122.5 0.0179
216 0.0172
384 0.0167
486 0.0168
0.0196

2.5 0.133
0.5 0.25
1.5 0.5
2.5 0.0263
3 0.0359
0.5 0.0501
4 0.0582
5 0.0582
6 0.0526
7 0.0537
8 0.0581
9 0.0675
10 0.0659
11 0.0669
12 0.0673
13 0.0669
14 0.0699
15 0.0745

995300
BPE/C18/.95EN/30C/L

995600
BPE/C18/.95EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
11.5	0.9254	0.0254	0.9254
12.3.28	0.0469	0.0478	0.0469
15.15	0.0668	0.0668	0.0668
17.5	0.0957	0.121	0.0957
23.7	0.263	0.213	0.263
30.1	0.335	0.313	0.335
43.15	0.506	0.506	0.506
53.	0.594	0.594	0.594
121	0.698	0.698	0.698
170	0.696	0.696	0.696
289	0.746	0.746	0.746
389	0.747	0.747	0.747
457	0.782	0.782	0.782
553	0.788	0.788	0.788
796	0.819	0.819	0.819
961	0.919	0.919	0.919
965			

940301
BPE/C18/OD/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0134	0.017	0.017
0.033	0.0198	0.033	0.033
0.067	0.0293	0.067	0.067
0.117	0.0395	0.117	0.117
0.183	0.0508	0.183	0.183
0.25	0.0602	0.25	0.25
0.5	0.0883	0.5	0.5
0.75	0.111	0.75	0.75
1.5	0.13	1.5	1.5
2.5	0.162	2.5	2.5
4.5	0.19	4.5	4.5
5.5	0.214	5.5	5.5
8	0.277	8	8
24	0.329	24	24
33.7	0.404	33.7	33.7
48	0.706	48	48
56.2	0.86	56.2	56.2
72.3	0.941	72.3	72.3
98.6	0.956	98.6	98.6
172	0.966	172	172
	0.972		0.972
	0.978		0.978

940601
BPE/C18/OD/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0134	0.017	0.017
0.033	0.0198	0.033	0.033
0.067	0.0293	0.067	0.067
0.117	0.0395	0.117	0.117
0.183	0.0508	0.183	0.183
0.25	0.0602	0.25	0.25
0.5	0.0883	0.5	0.5
0.75	0.111	0.75	0.75
1.5	0.13	1.5	1.5
2.5	0.162	2.5	2.5
4.5	0.19	4.5	4.5
5.5	0.214	5.5	5.5
8	0.277	8	8
24	0.329	24	24
33.7	0.404	33.7	33.7
48	0.706	48	48
56.2	0.86	56.2	56.2
72.3	0.941	72.3	72.3
98.6	0.956	98.6	98.6
172	0.966	172	172
	0.972		0.972
	0.978		0.978

BPE/C18/ON/300/L

BPE/C18/ON/600/L

t, h Mt/Mo

0.117	0.00905
0.183	0.0151
0.3	0.0212
0.5	0.0282
0.833	0.0427
1.5	0.0616
2.5	0.0826
4.25	0.107
7.03	0.151
22.8	0.297
220.9	0.325
48.1	0.425
123	0.658
217	0.788
386	0.881
626	0.913
889	0.927
1202	0.937
1494	0.895
2019	0.947

t, h Mt/Mo

0.017	0.0557
0.033	0.0717
0.067	0.0942
0.133	0.134
0.25	0.193
0.5	0.293
1	0.444
2	0.64
4	0.85
6	0.932
7.38	0.957
24	0.982
30.7	0.976
75.5	0.98
149	0.992

BPE/C18/TB/30C/L

960600
BPE/C18/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 133	0. 0157	0. 17	0. 0287
0. 25	0. 0245	0. 33	0. 0398
0. 5	0. 0264	0. 67	0. 0503
1. 98	0. 0456	1. 17	0. 0703
4	0. 0912	1.83	0. 103
5	0. 117	2.5	0. 11
6	0. 125	3.5	0. 265
7. 42	3.46	4.1	0. 314
23. 5	21.3	5.5	0. 491
57. 5	34.6	7.5	0. 47
95. 1	46.6	9.2	0. 532
145	54.4	11.2	0. 643
196	59.9	13.1	0. 761
264	66.7	15.5	0. 847
312	69.4	17.25	0. 918
361	72.5	19.2	0. 986
432	75.2	21.7	0. 992
484	77.7	24.7	0. 978
529	78.7	27.5	0. 983
601	81.3	30.5	0. 97
649	81.1	33.9	0. 988
769	83.2	38.9	0. 989
989	85.4		

980600
BPE/C18/AQ/60C/L

t, h

	Mt/Mo
3292	0.00434
4036	0.00727
4731	0.0112
5593	0.0218
6312	0.0311
7057	0.045
7540	0.059

BPE/C18/HB/40C/L

BPE/C18/HB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.5	0.0522	0.017	0.0273
1.1	0.0783	0.033	0.0238
1.97	0.103	0.067	0.074
4.4	0.172	0.133	0.108
5.53	0.235	0.25	0.15
6.78	0.254	0.5	0.246
24.3	0.502	1	0.364
30.5	0.561	2	0.531
54.7	0.664	3	0.646
145	0.722	20.5	0.957
193	0.805	24.1	0.976
313	0.848	25.7	0.974
409	0.867	44.8	0.987
526	0.846	141	0.984
724	0.869	1359	0.979

BPE/C18/HB/60C/L

BPE/C18/HB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.5	0.017	0.017	0.0273
1.1	0.033	0.033	0.0238
1.97	0.067	0.067	0.074
4.4	0.133	0.133	0.108
5.53	0.25	0.25	0.15
6.78	0.5	0.5	0.246
24.3	1	1	0.364
30.5	2	2	0.531
54.7	3	3	0.646
145	20.5	20.5	0.957
193	24.1	24.1	0.976
313	44.8	44.8	0.974
409	141	141	0.987
526	1359	1359	0.984
724	724	724	0.979

1120600
LPE/C18/EH/60C/L

t, h	Mt/Mo	LPE/C18/EH/60C/L	t, h	Mt/Mo	LPE/C18/EH/60C/L
0.017	0.00278	0.017	0.0019	0.00304	
0.033	0.00381	0.033	0.00304	0.00489	
0.067	0.00635	0.067	0.0076	0.0111	
0.133	0.00849	0.133	0.0111	0.017	
0.25	0.0126	0.25	0.0259	0.0451	
0.5	0.022	0.5	0.033	0.0607	
1.1	0.0326	1.1	0.0739	0.0939	
1.5	0.0397	1.5	0.187	0.227	
2.1	0.0506	2.1	0.42	0.45	
2.5	0.0506	2.5	0.55	0.55	
4.4	0.0637	4.4	0.62	0.618	
5.5	0.0732	5.5	0.67	0.736	
8.8	0.087	8.8	0.797	0.838	
19.2	0.141	19.2	0.861	0.861	
31.3	0.16	31.3	0.96	0.96	
95.9	0.265	95.9	0.993	0.993	
192	0.356	192	0.994	0.994	
364	0.486	364	0.998	0.998	
605	0.595	605	0.998	0.998	
990	0.69	990	0.998	0.998	
1200	0.724	1200	0.998	0.998	
1612	0.803	1612	0.998	0.998	
1613	0.802	1613	0.998	0.998	
1872	0.822	1872	0.998	0.998	
2094	0.775	2094	0.998	0.998	
2862	0.746	2862	0.998	0.998	
3652	0.869	3652	0.998	0.998	
4999	0.858	4999	0.998	0.998	
5715	0.907	5715	0.998	0.998	

1520600
 BPE/C32/EH/60C/L
 t, h Mt/Mo

0.017	0.0109
0.033	0.0154
0.067	0.0201
0.133	0.0287
0.25	0.0441
0.5	0.0693
1	0.104
1.5	0.13
2.5	0.171
4	0.218
5.5	0.254
8	0.303
24	0.404
55.3	0.625
121	0.693
289	0.725
384	0.74
676	0.749
677	0.75
865	0.751
1183	0.757
1538	0.76
1997	0.755
2191	

1530300
 BPE/C32/HP/30C/L
 t, h Mt/Mo

0.017	0.017
0.033	0.033
0.067	0.067
0.133	0.133
0.25	0.121
0.5	0.198
1	0.322
1.5	0.528
2.5	0.935
4	1.21
6	1.36
9	1.67
14	3.07
23	3.79
30	4.34
47	4.79
67	5.25
145	7.15
241	7.32
508	7.41
697	7.55
894	7.59
1153	7.51
1489	7.83
1732	8

1530601
 BPE/C32/HP/60C/U
 t, h Mt/Mo

0.017	0.017
0.033	0.033
0.067	0.067
0.133	0.133
0.25	0.25
0.5	0.5
1	0.558
1.5	0.888
2.5	0.94
4	0.959
6	0.965
9	0.971
14	0.984
24	0.987
47	0.984
67	0.987
145	0.992
241	0.992
508	0.992
697	0.993
894	0.994
1153	0.994
1489	0.994
1732	0.994
2043	0.994

1610300
BPE/C32/C0/30C/L

1610600
BPE/C32/C0/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	9.35E-4	0.017	0.0132
0.033	0.00106	0.033	0.0194
0.067	0.00164	0.067	0.0282
0.133	0.00226	0.133	0.0381
0.2	0.0028	0.2	0.0529
0.25	0.00342	0.25	0.0618
0.5	0.00489	0.75	0.11
1.5	0.00654	1.5	0.154
2.5	0.00789	2.5	0.176
5.5	0.0118	2.5	0.242
11.5	0.0135	4.5	0.29
22.5	0.0175	7.83	0.33
45.5	0.0209	24.3	0.433
91.5	0.0271	31.8	0.508
182.5	0.0437	97.8	0.595
365.5	0.0504	194	0.811
732.5	0.0604	360	0.85
147.8	0.0779	602	0.912
295.8	0.103	844	0.952
591.8	0.123	1327	0.947
1183.8	0.155		0.958
2367.8	0.185		
4735.8	0.219		
9471.8	0.25		
18943.8	0.29		
37887.8	0.306		
75774.8	0.339		
15548.8	0.364		
31092.8	0.335		
62144.8	0.405		
124148.8	0.421		
24887.8	0.436		
49750.8	0.45		
99500.8	0.462		
19902.8	0.473		
39894.8	0.482		

1620300
BPE/C32/EH/30C/L1620600
BPE/C32/EH/60C/L

t, h	Nt/Mo	t, h	Nt/Mo
0.017	2. 41E-4	0.017	0. 002
0.033	3. 19E-4	0.033	0. 0263
0.067	4. 95E-4	0.067	0. 0341
0.133	8. 26E-4	0.133	0. 0515
0.25	0. 00132	0.25	0. 0815
0.5	0. 00217	0.5	0. 138
1	0. 00347	1	0. 217
1.5	0. 00469	1.5	0. 275
2.5	0. 00649	2.5	0. 363
4.07	0. 0093	4	0. 462
5.5	0. 0114	5	0. 622
7.8	0. 0146	7.8	0. 803
23.7	0. 0277	24.3	0. 837
31.8	0. 0334	31.9	0. 835
55.8	0. 0437	55.9	0. 835
104	0. 059	79.7	0. 848
264	0. 0981	151	0. 848
432	0. 100	245	0. 849
654	0. 126	394	0. 844
1422	0. 145	486	0. 854
2211	0. 166	700	0. 851
3559	0. 166	988	0. 856
4638	0. 152	1248	0. 858
5310	0. 145	1470	0. 847
6389	0. 156	1710	0. 857

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
1625600 BPE/C32/.5EN/60C/L	1695300 BPE/C32/.95EN/30C/L	1695600 BPE/C32/.95EN/60C/L			
0. 017	2. 55E-4	0. 033	2. 08E-4		
0. 033	3. 78E-4	0. 067	3. 26E-4		
0. 067	5. 06E-4	0. 133	5. 08E-4		
0. 133	7. 51E-4	0. 25	8. 6E-4		
0. 25	0. 5	0. 5	0. 00137		
0. 5	0. 00197	1	0. 00231		
1	0. 00323	2. 05	0. 00391		
1. 5	0. 00421	4	0. 00635		
2. 5	0. 00561	5. 37	0. 00787		
4. 5	0. 00847	23. ?	0. 0203		
7. 8?	0. 00961	29. 5	0. 0238		
23. 7	0. 0114	48. 5	0. 0319		
31. 9	0. 0121	77. 2	0. 0403		
55. 9	0. 0123	149	0. 0552		
79. 5	0. 0122	316	0. 0754		
240	0. 0125	504	0. 0923		
408	0. 0109	673	0. 0991		
630	0. 0103	841	0. 108		
2697	0. 0107	1177	0. 117		
		1441	0. 116		
		1733	0. 123		
		1901	0. 137		

1630301 BPE/C32/HP/30C/U
 t, h Mt/Mo
 0.017 0.00788
 0.033 0.012
 0.067 0.0178
 0.0133 0.0272
 0.025 0.0361
 0.025 0.0422
 0.025 0.0742
 0.025 0.104
 0.025 0.128
 0.025 0.158
 0.025 0.178
 0.025 0.198
 0.025 0.218
 0.025 0.238
 0.025 0.258
 0.025 0.278
 0.025 0.298
 0.025 0.318
 0.025 0.338
 0.025 0.358
 0.025 0.378
 0.025 0.398
 0.025 0.418
 0.025 0.438
 0.025 0.458
 0.025 0.478
 0.025 0.498
 0.025 0.518
 0.025 0.538
 0.025 0.558
 0.025 0.578
 0.025 0.598
 0.025 0.618
 0.025 0.638
 0.025 0.658
 0.025 0.678
 0.025 0.698
 0.025 0.718
 0.025 0.738
 0.025 0.758
 0.025 0.778
 0.025 0.798
 0.025 0.818
 0.025 0.838
 0.025 0.858
 0.025 0.878
 0.025 0.898
 0.025 0.918
 0.025 0.938
 0.025 0.958
 0.025 0.978
 0.025 0.998

1630601 BPE/C32/HP/60C/U
 t, h Mt/Mo
 0.017 0.016
 0.033 0.251
 0.067 0.401
 0.133 0.662
 0.2 0.791
 0.25 0.841
 0.5 0.916
 0.75 0.934
 1 0.942
 1.5 0.955
 2 0.965
 2.5 0.972
 5 0.978
 5 0.982
 5 0.985
 5 0.988
 5 0.991
 5 0.994
 5 0.997
 5 0.999
 8 0.982
 8 0.985
 8 0.988
 8 0.991
 8 0.994
 8 0.997
 8 0.999
 12.3 0.982
 12.3 0.985
 12.3 0.988
 12.3 0.991
 12.3 0.994
 12.3 0.997
 12.3 0.999
 15.65 0.991
 15.65 0.994
 15.65 0.997
 15.65 0.999
 16.61 0.991
 16.61 0.994
 16.61 0.997
 16.61 0.999

1650300
BPE/C32/0H/30C/L

1650600
BPE/C32/0H/60C/L

t, h	Nt/Mo	t, h	Nt/Mo
0. 017	1. 7E-4	0. 017	0. 0296
0. 033	2. 52E-4	0. 033	0. 0371
0. 067	3. 79E-4	0. 067	0. 0471
0. 117	6. 08E-4	0. 133	0. 0654
0. 183	9. 4E-4	0. 25	0. 103
0. 3	0. 00136	0. 5	0. 169
0. 5	0. 00207	1. 2	0. 262
0. 833	0. 00309	2. 4	0. 397
1. 5	0. 005	6	0. 572
2. 5	0. 00748	7.	0. 744
4. 25	0. 0112	23.	0. 874
7	0. 0164	31.	0. 891
24	0. 0384	48.	0. 893
51.	0. 0464	99.	0. 924
98.	0. 0645	145.	0. 919
200	0. 0969	217	0. 936
362	0. 151		
602	0. 23		
865	0. 284		
1178	0. 364		
1470	0. 395		
1897	0. 428		
2401	0. 459		
2906	0. 483		
3488	0. 505		
4081	0. 525		
4729	0. 543		
5435	0. 558		
6169	0. 574		
6264	0. 581		

1660300
BPE/C32/TB/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	3.21E-4	0.017	0.019
0.033	5.33E-4	0.033	0.027
0.067	9.16E-4	0.067	0.0387
0.117	0.0014	0.133	0.0558
0.183	0.00204	0.25	0.0814
0.3	0.00297	0.5	0.13
0.5	0.0044	1	0.199
0.833	0.00646	2	0.291
1.5	0.0101	4	0.426
2.5	0.0145	6	0.52
3.3	0.0199	9	0.581
5.3	0.0253	23	0.805
7.25	0.0308	30	0.832
23	0.0618	48.	0.865
31.3	0.0746	72	0.877
51.9	0.0976	169	0.906
79.3	0.123	240	0.905
144	0.168	340	0.915
220	0.208	505	0.921
385	0.277	673	0.916
579	0.342	961	0.932
871	0.405		
1201	0.464		
1538	0.514		
1925	0.563		
2401	0.613		
2931	0.659		
3412	0.701		

1660600
BPE/C32/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	0.017	0.019
0.033	0.033	0.067	0.0387
0.067	0.067	0.133	0.0558
0.117	0.133	0.25	0.0814
0.183	0.25	0.5	0.13
0.3	0.5	1	0.199
0.5	2	2	0.291
0.833	4	4	0.426
1.5	6	6	0.52
2.5	9	9	0.581
3.3	23	30	0.805
5.3	30	48.	0.832
7.25	48.	72	0.865
23	72	169	0.877
31.3	169	240	0.906
51.9	240	340	0.905
79.3	340	505	0.915
144	505	673	0.921
220	673	961	0.932
385	961		
579			
871			
1201			
1538			
1925			
2401			
2931			
3412			

1670300
BPE/C32/T0/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	2.7E-4	0.017	0.0106
0.033	4.17E-4	0.033	0.0157
0.067	6.34E-4	0.067	0.0231
0.133	9.36E-4	0.133	0.0417
0.25	0.00128	0.25	0.0609
0.5	0.00148	0.5	0.0735
1.5	0.00243	1.5	0.136
3.5	0.00337	3.5	0.181
7.5	0.00424	7.5	0.218
15.	0.00569	15.	0.281
35.	0.00703	35.	0.333
75.	0.00825	75.	0.379
155.	0.0115	155.	0.429
355.	0.0142	355.	0.574
755.	0.0184	755.	0.658
1555.	0.0361	1555.	0.846
3555.	0.0552	3555.	0.874
7555.	0.0749	7555.	0.896
15555.	0.102	15555.	0.905
35555.	0.123	35555.	0.919
75555.	0.16	75555.	0.937
155555.	0.197	155555.	0.947
355555.	0.239	355555.	0.948
755555.	0.271	755555.	0.951
1555555.	0.302	1555555.	0.948
3555555.	0.343	3555555.	0.954
7555555.	0.358	7555555.	1.688
15555555.	0.389		
35555555.	0.408		
75555555.	0.425		
155555555.	0.44		
355555555.	0.452		
755555555.	0.47		
1555555555.	0.47		

60.7

1680600
BPE-C32/HQ/68C/L

t, h

124.8

24.4

48.4

122

725

1346

397

725

11925

119297

33269

4012

4708

5593

6289

7034

7538

1. 81E-4

2. 55E-4

3. 42E-4

4. 06E-4

5. 6. 83E-4

7. 84E-4

8. 00132

9. 0016

0. 00225

0. 00275

0. 00303

0. 00377

0. 00417

0. 00563

0. 00793

0. 0102

0. 0125

0. 0175

1690400
BPE/C32/HB/40C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	8.12E-4	0.017	0.0298
0.033	0.00231	0.033	0.0328
0.067	0.00255	0.067	0.0497
0.15	0.00614	0.133	0.0662
0.25	0.00954	0.25	0.0991
0.5	0.016	0.5	0.167
1	0.025	1	0.248
1.97	0.0378	1.95	0.35
4	0.0562	4	0.516
6	0.0705	6	0.856
7.42	0.0783	7.5	0.882
24.3	0.14	23.5	0.895
30.3	0.158	30.6	0.919
51.9	0.205	51.4	0.936
121	0.307	129	0.943
169	0.36	133	0.947
221	0.409	481	0.954
289	0.458	508	
385	0.515		
556	0.586		
864	0.651		
1009	0.674		
1301	0.698		
1468	0.71		
1732	0.725		
1993	0.736		
2329	0.751		
2405	0.753		

LPE/C32/C0/30C/L	t, h	Mt/Mo
0.017	1.77E-4	
0.033	1.52E-4	
0.067	1.94E-4	
0.133	2.97E-4	
0.25	3.26E-4	
0.5	6.05E-4	
1	0.00105	
1.5	0.00143	
2.5	0.00197	
4.5	0.00273	
8.5	0.00348	
17	0.00459	
34	0.00568	
68	0.00733	
137	0.0119	
274	0.0155	
547	0.0207	
1094	0.0272	
2188	0.0324	
4377	0.0421	
8754	0.0498	
17518	0.0584	
35036	0.0643	
70072	0.071	
140144	0.0762	
280288	0.084	
560576	0.0898	
1121152	0.0943	
2242304	0.0966	
4484608	0.1036	
8969216	0.1073	
17938432	0.112	

LPE/C32/C0/60C/L	t, h	Mt/Mo
0.017	0.017	
0.033	0.033	
0.067	0.067	
0.133	0.133	
0.25	0.25	
0.5	0.5	
1	0.75	
1.5	1.15	
2.5	1.5	
4.5	2.5	
8.5	5	
17	5	
34	5	
68	5	
137	5	
274	5	
547	5	
1094	5	
2188	5	
4377	5	
8754	5	
17518	5	
35036	5	
70072	5	
140144	5	
280288	5	
560576	5	
1121152	5	
2242304	5	
4484608	5	
8969216	5	
17938432	5	

1920300
LPE/C32/EH/30C/L

t, h Mt/Mo
LPE/C32/EH/60C/L

0.067	1.75E-4	0.017	0.00324
0.133	2.94E-4	0.033	0.00476
0.25	4.02E-4	0.067	0.00705
0.5	7.69E-4	0.00122	0.0219
1.2	0.00213	0.0047	0.0365
2.4	0.00349	0.0056	0.0558
4.6	0.0047	0.0112	0.0741
7.6	0.0056	0.0125	0.0955
12.4	0.0123	0.0223	0.112
24.1	0.0262	0.0301	0.125
48.	0.0325	0.031	0.138
72.	0.0325	0.035	0.155
149	0.0331	0.035	0.173
245	0.0353	0.035	0.187
409	0.0353	0.035	0.207
580	0.0362	0.036	0.228
721	0.0362	0.036	0.248
846	0.0351	0.035	0.267
1061	0.0351	0.035	0.284
1228	0.0351	0.035	0.303
1395	0.0351	0.035	0.323
1657	0.0351	0.035	0.343
1877	0.0351	0.035	0.363
2140	0.0375	0.0383	0.383
2581	0.0383	0.0385	0.403
3365	0.0385	0.0385	0.423
3365	0.0385	0.0385	0.443

1925600
LPE/C32/EH/60C/L

t, h Mt/Mo
LPE/C32/EH/60C/L

0.5	1.59E-4	0.1	1.5
1.1	3.51E-4	2.5	2.5
2.5	3.52E-4	4.1	4.1
5.5	7.02E-4	7.85	7.85
11.0	0.00135	18.78	18.78
21.6	0.00167	23.84	23.84
43.2	0.00262	55.9	55.9
86.2	0.00396	111.0	111.0

1930301
LPE/C32/HP/30C/U

1930601
LPE/C32/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00242	0.017	0.0282
0.033	0.00398	0.033	0.0414
0.067	0.00656	0.067	0.0611
0.133	0.0102	0.133	0.0908
0.25	0.0149	0.25	0.132
0.5	0.022	0.5	0.205
1.2	0.0323	1.2	0.332
4.6	0.0489	4.6	0.567
7.5	0.0829	7.5	0.785
24.1	0.115	24.1	0.889
31.5	0.134	31.5	0.909
35.4	0.245	35.4	0.924
77.9	0.274	77.9	0.936
172	0.332	172	0.941
223	0.455	223	0.947
313	0.483	313	0.952
387	0.519	387	0.954
416	0.541	416	0.956
583	0.549	583	0.956
747	0.602	747	0.956
917	0.618	917	0.956
1057	0.628	1057	0.956
1326	0.638	1326	0.956
1540	0.649	1540	0.956
1708	0.659	1708	0.956
1875	0.668	1875	0.956
2137	0.678	2137	0.956
2357	0.686	2357	0.956
2620	0.694	2620	0.956
3002	0.704	3002	0.956
3265	0.712	3265	0.956
4243	0.722	4243	0.956
4823	0.734	4823	0.956

LPE/C32/DH/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.0.3	2.42E-4	7.1E-4	0.00119
0.0.5	0.833	0.00197	0.00301
1.5	2.5	0.00301	0.00438
4.25	7	0.00634	0.0159
24	31.1	0.0137	0.0207
79	49.1	0.027	0.0438
148	1241	0.0517	0.0616
384	576	0.0716	0.0845
576	869	0.0959	0.107
1182	11541	0.0959	0.116
1944	1541	0.116	0.125
2400	2400	0.125	0.134
2932	2932	0.134	0.14
3460	4079	0.14	0.15
4732	4732	0.15	0.16
5430	6101	0.16	0.171

LPE/C32/0H/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00321	0.033	0.0046
0.067	0.00718	0.133	0.0116
0.25	0.0195	0.5	0.0317
1.97	1.03	4	0.05
6.68	6.98	24.3	249.4
121	168	222	345
121	168	221	345
1543	1543	457	491
637	637	636	637
718	718	756	756
764	764	764	764
746	746	744	744
787	787	787	787
795	795	1830	1830
789	789	2185	2185
804	804	2669	2669

1970300
LPE/C32/T0/30C/L

t, h	Mt/Mo	
0. 017	2.14E-4	
0. 033	1.53E-4	
0. 067	1.99E-4	
0. 133	4.19E-4	
0. 25	5.64E-4	
0. 5	8.6E-4	
1. 75	0.0126	
2. 25	0.0153	
5. 2	0.0221	
11. 5	0.0263	
22. 5	0.0321	
45. 13	0.0418	
88. 13	0.0577	
179. 8	0.0798	
322. 1	0.1196	
483. 3	0.235	
791. 8	0.291	
146	0.372	
224	0.433	
389	0.547	
576	0.588	
844	0.676	
1160	0.6739	
1537	0.6803	
1904	0.6845	
2413	0.6925	
2887	0.6979	
3363	0.702	
3870	0.713	
4346	0.714	
4879	0.719	
5384	0.721	

1970600
LPE/C32/T0/60C/L

t, h	Mt/Mo	
0. 017	0.00239	
0. 033	0.00414	
0. 067	0.00633	
0. 133	0.0112	
0. 25	0.0147	
0. 5	0.0185	
1. 25	0.0315	
2. 5	0.0405	
5. 5	0.059	
11. 5	0.0672	
23. 5	0.073	
47. 5	0.0891	
95. 5	0.0993	
194	0.116	
392	0.174	
784	0.226	
1568	0.242	
3136	0.332	
6272	0.427	
12544	0.561	
25088	0.63	
50176	0.69	
100352	0.713	
200704	0.736	
401408	0.762	
802816	0.794	
1605632	0.8	
3211264	0.811	
6422528	0.816	
12845056	0.819	

LPE/C32/AQ/60C/L

t, h Mt/Mo

5.3
23.
119
316
456
623
816
1205
1611
1775
2111
2495
2783
3119
3627
4034
4703
3.25E-6
1.04E-4
2.21E-4
7.09E-4
0.00192
0.0016
0.00212
0.00418
0.00649
0.00782
0.0104
0.0143
0.0166
0.0197
0.0241
0.0272
0.0314

2010300
LPE/BHT/CO/30C/L2010600
LPE/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.067	2.85E-4	0.017	6.86E-4
0.117	4.27E-4	0.033	0.00111
0.183	3.95E-4	0.067	0.00148
0.25	5.21E-4	0.117	0.00269
0.5	9.33E-4	0.183	0.00421
0.75	1.0E-3	0.025	0.0059
1.5	0.00143	0.05	0.0119
2.5	0.00184	0.075	0.0181
4.03	0.00245	0.115	0.0224
5.5	0.00286	0.15	0.0305
8	0.00306	0.181	0.0371
24	0.00326	0.222	0.0427
32	0.00366	0.24	0.0457
48.4	0.00418	0.28	0.0688
56.2	0.00468	0.32	0.0825
120	0.00528	0.37	0.145
192	0.00662	0.47	0.166
364	0.00991	0.56	0.208
605	0.115	0.68	0.226
841	0.135	0.74	0.325
1158	0.156	0.8	0.422
1547	0.176	0.9	0.503
1886	0.194	0.96	0.538
2331	0.213	1.02	0.584
2858	0.234	1.14	0.595

2020300
LPE/BHT/EN/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	3.3E-4	0.017	0.00109
0.033	4.36E-4	0.033	0.00145
0.067	5.25E-4	0.067	0.00217
0.133	7.05E-4	0.133	0.00279
0.25	8.6E-4	0.25	0.00524
0.5	0.00126	0.5	0.00925
1.1	0.00179	1.1	0.0168
1.5	0.00222	1.2	0.0266
2.5	0.00347	2.4	0.0421
4.4	0.00462	6.5	0.0505
23.6	0.00592	23.8	0.0601
31.5	0.00767	31.5	0.111
100	0.0156	127	0.126
192	0.0187	127	0.166
360	0.0365	53.5	0.24
915	0.0519	220	0.297
1495	0.0708	389	0.359
1755	0.107	460	0.394
2197	0.129	733	0.444
3027	0.137	1060	0.455
3027	0.151	1705	0.493
	0.185	1705	0.495
	0.188		

2020600
LPE/BHT/EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	0.017	0.00109
0.033	0.033	0.033	0.00145
0.067	0.067	0.067	0.00217
0.133	0.133	0.133	0.00279
0.25	0.25	0.25	0.00524
0.5	0.5	0.5	0.00925
1.1	1.1	1.1	0.0168
1.5	1.2	1.2	0.0266
2.5	2.4	2.4	0.0421
4.4	6.5	6.5	0.0505
23.8	23.1	23.1	0.0601
31.5	31.5	31.5	0.111
100	127	127	0.126
192	53.5	53.5	0.166
360	220	220	0.24
915	389	389	0.297
1495	460	460	0.359
1755	733	733	0.394
2197	1060	1060	0.444
3027	1705	1705	0.455
3027	1705	1705	0.493
	1705	1705	0.495

2030301 LPE/BHT/HP/30C/U

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0. 017	0.00372	0. 017	0.0201	0. 017	0.0201	0. 017	0.0201
0. 033	0.00676	0. 033	0.0322	0. 067	0.0488	0. 067	0.0488
0. 067	0.0124	0. 067	0.0754	0. 133	0.133	0. 133	0.133
0. 133	0.0207	0. 133	0.0968	0. 225	0.225	0. 225	0.225
0. 2	0.027	0. 25	0.111	1. 5	1.5	1. 5	1.5
0. 25	0.0314	0. 5	0.177	2. 25	2.25	2. 25	2.25
0. 5	0.0495	0. 75	0.24	3. 25	3.25	3. 25	3.25
0. 783	0.0677	1.	0.304	4. 25	4.25	4. 25	4.25
1	0.091	1. 5	0.431	5. 25	5.25	5. 25	5.25
1. 5	0.111	2	0.504	6. 25	6.25	6. 25	6.25
2. 02	0.141	2. 5	0.541	7. 25	7.25	7. 25	7.25
2. 5	0.17	4. 5	0.58	8. 25	8.25	8. 25	8.25
4. 5	0.269	6. 5	0.62	9. 25	9.25	9. 25	9.25
5. 5	0.378	8. 5	0.66	10. 25	10.25	10. 25	10.25
8	0.499	11	0.71	11. 25	11.25	11. 25	11.25
11	0.602	13	0.82	13. 25	13.25	13. 25	13.25
13	0.607	15	0.93	15. 25	15.25	15. 25	15.25
15	0.611	17	1.04	17. 25	17.25	17. 25	17.25
17	0.614	19	1.15	19. 25	19.25	19. 25	19.25
19	0.617	21	1.26	21. 25	21.25	21. 25	21.25
21	0.617	23	1.37	23. 25	23.25	23. 25	23.25
23	0.618	25	1.48	25. 25	25.25	25. 25	25.25
25	0.619	27	1.59	27. 25	27.25	27. 25	27.25
27	0.620	29	1.70	29. 25	29.25	29. 25	29.25
29	0.621	31	1.81	31. 25	31.25	31. 25	31.25
31	0.622	33	1.92	33. 25	33.25	33. 25	33.25
33	0.623	35	2.03	35. 25	35.25	35. 25	35.25
35	0.624	37	2.14	37. 25	37.25	37. 25	37.25
37	0.625	39	2.25	39. 25	39.25	39. 25	39.25
39	0.626	41	2.36	41. 25	41.25	41. 25	41.25
41	0.627	43	2.47	43. 25	43.25	43. 25	43.25
43	0.628	45	2.58	45. 25	45.25	45. 25	45.25
45	0.629	47	2.69	47. 25	47.25	47. 25	47.25
47	0.630	49	2.80	49. 25	49.25	49. 25	49.25
49	0.631	51	2.91	51. 25	51.25	51. 25	51.25
51	0.632	53	3.02	53. 25	53.25	53. 25	53.25
53	0.633	55	3.13	55. 25	55.25	55. 25	55.25
55	0.634	57	3.24	57. 25	57.25	57. 25	57.25
57	0.635	59	3.35	59. 25	59.25	59. 25	59.25
59	0.636	61	3.46	61. 25	61.25	61. 25	61.25
61	0.637	63	3.57	63. 25	63.25	63. 25	63.25
63	0.638	65	3.68	65. 25	65.25	65. 25	65.25
65	0.639	67	3.79	67. 25	67.25	67. 25	67.25
67	0.640	69	3.90	69. 25	69.25	69. 25	69.25
69	0.641	71	4.01	71. 25	71.25	71. 25	71.25
71	0.642	73	4.12	73. 25	73.25	73. 25	73.25
73	0.643	75	4.23	75. 25	75.25	75. 25	75.25
75	0.644	77	4.34	77. 25	77.25	77. 25	77.25
77	0.645	79	4.45	79. 25	79.25	79. 25	79.25
79	0.646	81	4.56	81. 25	81.25	81. 25	81.25
81	0.647	83	4.67	83. 25	83.25	83. 25	83.25
83	0.648	85	4.78	85. 25	85.25	85. 25	85.25
85	0.649	87	4.89	87. 25	87.25	87. 25	87.25
87	0.650	89	5.00	89. 25	89.25	89. 25	89.25
89	0.651	91	5.11	91. 25	91.25	91. 25	91.25

2030602 LPE/BHT/HP/60C/U2

t, h	Mt/Mo						
0. 017	0.017	0. 017	0.017	0. 017	0.017	0. 017	0.017
0. 033	0.033	0. 033	0.033	0. 067	0.067	0. 067	0.067
0. 067	0.067	0. 133	0.133	0. 133	0.133	0. 133	0.133
0. 133	0.133	0. 227	0.227	0. 227	0.227	0. 227	0.227
0. 227	0.227	0. 314	0.314	0. 314	0.314	0. 314	0.314
0. 314	0.314	0. 401	0.401	0. 401	0.401	0. 401	0.401
0. 401	0.401	0. 488	0.488	0. 488	0.488	0. 488	0.488
0. 488	0.488	0. 574	0.574	0. 574	0.574	0. 574	0.574
0. 574	0.574	0. 660	0.660	0. 660	0.660	0. 660	0.660
0. 660	0.660	0. 746	0.746	0. 746	0.746	0. 746	0.746
0. 746	0.746	0. 832	0.832	0. 832	0.832	0. 832	0.832
0. 832	0.832	0. 918	0.918	0. 918	0.918	0. 918	0.918

2070300
LPE/BHT/T0/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	3.09E-4	0.017	7.13E-4
0.033	3.71E-4	0.033	0.00102
0.067	4.7E-4	0.067	0.00169
0.117	4.78E-4	0.117	0.00229
0.183	5.17E-4	0.0:183	0.00368
0.25	7.3E-4	0.0:25	0.0046
0.5	7.89E-4	0.0:75	0.00976
1.5	0.00139	1.5	0.0143
2.5	0.00164	2.5	0.0244
4	0.00202	4	0.0305
5	0.00243	5	0.0354
8	0.00366	8	0.0487
12	0.00461	12	0.0605
24	0.0148	24	0.0761
32	0.0183	32	0.0761
48	0.0246	48	0.145
76.	0.0322	76.	0.172
128	0.0469	128	0.214
221	0.0655	221	0.311
386	0.0943	386	0.375
631	0.122	631	0.434
899	0.151	899	0.536
1238	0.179	1238	0.597
1566	0.205	1566	0.602
1971	0.23	1971	0.607
2214	0.245		

2130601
LPE/BHT/HP/60C/U

t, h	Mt/Mo
0.017	0.0925
0.033	0.139
0.067	0.212
0.133	0.335
0.25	0.531
0.5	0.648
1	0.666
2	0.668
4.5	0.669
7	0.67
24.3	0.671

2210300
LPE/BHT/CO/38C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	5.25E-4	0.017	0.00443
0.033	8.6E-4	0.033	0.00729
0.067	0.00112	0.067	0.0124
0.133	0.00181	0.133	0.0186
0.25	0.00299	0.25	0.0317
0.5	0.00569	0.5	0.0577
1.1.5	0.0105	1.1.5	0.0917
2.5	0.0221	2.5	0.117
4.5.5	0.0303	4.5.5	0.158
7.58	0.0379	8.03	0.202
24.3	0.0444	24.1	0.239
32.2	0.0513	32.1	0.294
52.5	0.0796	50.1	0.313
96.4	0.0913	97.7	0.399
193	0.115	193	0.604
362	0.156	362	0.604
607	0.225	607	0.743
866	0.304	866	0.909
1175	0.394	1175	0.952
1536	0.47	1536	0.954
1922	0.627	1922	0.964
2282	0.698	2282	0.964
2886	0.766	2886	0.968
3623	0.829	3623	0.891
4302	0.885		

2210600
LPE/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	0.017	0.00443
0.033	0.033	0.033	0.00729
0.067	0.067	0.067	0.0124
0.133	0.133	0.133	0.0186
0.25	0.25	0.25	0.0317
0.5	0.5	0.5	0.0577
1.1.5	1.1.5	1.1.5	0.0917
2.5	2.5	2.5	0.117
4.5.5	4.5.5	4.5.5	0.158
7.58	7.58	8.03	0.202
24.1	24.1	24.1	0.239
32.1	32.1	32.1	0.294
50.1	50.1	50.1	0.313
97.7	97.7	97.7	0.399
193	193	193	0.604
362	362	362	0.604
607	607	607	0.743
866	866	866	0.909
1175	1175	1175	0.952
1536	1536	1536	0.954
1922	1922	1922	0.964
2282	2282	2282	0.964
2886	2886	2886	0.968
3623	3623	3623	0.968
4302	4302		

LPE/BHT/EN/30C/L

2220300
LPE/BHT/EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00102	0.017	0.0101	0.033	0.0148
0.033	0.00183	0.033	0.024	0.067	0.0379
0.067	0.00191	0.067	0.0589	0.133	0.133
0.133	0.00343	0.133	0.25	0.25	0.5
0.25	0.00506	0.25	1	0.517	2
0.517	0.09854	0.517	2	1.2	4
1.2	0.0226	1.2	4	2.4	6
2.4	0.0365	2.4	6	3.6	8
3.6	0.0503	3.6	8	5.6	12
5.6	0.059	5.6	12	7.5	23.8
7.5	0.103	7.5	23.8	9.4	48.9
9.4	0.139	9.4	48.9	12.4	72
12.4	0.174	12.4	72	19.3	193
19.3	0.241	19.3	193	38.5	385
38.5	0.319	38.5	385	60.1	601
60.1	0.391	60.1	601	75.0	750
75.0	0.433	75.0	750	91.3	913
91.3	0.473	91.3	913	105.7	1057
105.7	0.513	105.7	1057	122.8	1228
122.8	0.545	122.8	1228	134.5	1345
134.5	0.574	134.5	1345	153.7	1537
153.7	0.604	153.7	1537	170.8	1708
170.8	0.626	170.8	1708	189.7	1897
189.7	0.659	189.7	1897	208.9	2089
208.9	0.683	208.9	2089	240.5	2405
240.5	0.714	240.5	2405	271.3	2713
271.3	0.741	271.3	2713	300.4	3004
300.4	0.768	300.4	3004	336.0	3360
336.0	0.789	336.0	3360	405.6	4056
405.6	0.827	405.6	4056	475.6	4756
475.6	0.848	475.6	4756		

2230300
LPE/BHT/HP/30C/L

2230301
LPE/BHT/HP/30C/U

2230601
LPE/BHT/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0156	0.017	0.0306	0.033	0.0508	0.033	0.128
0.033	0.0281	0.033	0.067	0.067	0.0822	0.067	0.119
0.067	0.045	0.117	0.119	0.117	0.162	0.133	0.287
0.117	0.0687	0.183	0.183	0.25	0.25	0.25	0.439
0.183	0.0984	0.25	0.25	0.5	0.5	0.5	0.704
0.25	0.126	0.5	0.5	1.5	1.5	1.5	0.927
0.33	0.16	0.75	0.75	2.5	2.5	2.5	1.03
0.417	0.196	1.5	1.5	4.5	4.5	4.5	3.75
0.5	0.232	1.5	1.5	5.5	5.5	5.5	2.4
0.667	0.311	1.5	1.5	8.5	8.5	8.5	0.962
0.833	0.402	1.5	1.5	11.5	11.5	11.5	0.963
1.17	0.506	1.5	1.5	14.5	14.5	14.5	0.964
1.33	0.693	1.5	1.5	17.5	17.5	17.5	0.965
1.75	0.817	1.5	1.5	21.5	21.5	21.5	0.963
2.25	0.889	1.5	1.5	23.5	23.5	23.5	0.964
2.55	0.908	1.5	1.5	24.5	24.5	24.5	0.964
3.55	0.94	1.5	1.5	26.5	26.5	26.5	0.959
5.5	0.952	1.5	1.5	27.5	27.5	27.5	0.957
12.6	0.96	1.5	1.5	12.6	12.6	12.6	0.959
24	0.959	1.5	1.5	24	24	24	0.957
97.1	0.957	1.5	1.5	97.1	97.1	97.1	0.957

2250300
LPE/BHT/ON/30C/L

2250600
LPE/BHT/ON/60C/L1

2250602
LPE/BHT/ON/60C/L2

t, h Mt/Mo t, h Mt/Mo

t, h Mt/Mo t, h Mt/Mo

t, h Mt/Mo t, h Mt/Mo

0.067	0.0011	0.067	0.0282	0.017	0.00322
0.117	0.0017	0.133	0.0391	0.033	0.00507
0.183	0.00276	0.25	0.0569	0.067	0.00722
0.3	0.00405	0.5	0.0959	0.117	0.0109
0.5	0.00699	1	0.131	0.0	0.0161
0.85	0.0115	2	0.203	0.0	0.0254
1.45	0.0167	4	0.322	0.0	0.0434
2.45	0.0264	7	0.473	0.0	0.0716
6.65	0.0393	25.8	0.879	0.0	0.119
24.6	0.0507	28.9	0.893	0.0	0.185
30.6	0.124	42.7	0.923	0.1	0.275
54.5	0.143	82.4	0.928	0.5	0.372
124	0.222	120	0.927	33.6	0.492
1216	0.498	193	0.912	56	0.865
2784	0.725	289	0.944	120	0.893
624	0.854			217	0.914
865	0.835			385	0.931
1179	0.904				0.932
1599	0.918				

2260300
LPE/BHT/TB/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.133	0.00181	0.017	0.00616
0.25	0.00216	0.033	0.00737
0.5	0.00356	0.067	0.0136
1.1	0.00487	0.117	0.0197
2.2	0.00805	0.183	0.0272
4.4	0.0113	0.25	0.035
5.5	0.0132	0.5	0.0618
7.7	0.0153	1.5	0.0797
14.3	0.0319	2.5	0.094
23.4	0.0378	3.5	0.118
25.5	0.0518	5.5	0.137
51.7	0.0653	8.5	0.155
72.5	0.112	11.5	0.185
169	0.166	18.3	0.239
316	0.196	24.4	0.529
410	0.25	32.4	0.617
576	0.289	48.8	0.744
724	0.342	80.5	0.867
889	0.434	147	0.923
1255	0.531	225	0.931
1661	0.591	398	0.937
1901	0.62	577	0.931
2161	0.675	702	0.942
2545	0.733		
3169	0.753		
3365			

2260600
LPE/BHT/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00616	0.033	0.00737
0.067	0.0136	0.117	0.0197
0.183	0.0272	0.25	0.035
0.5	0.0618	0.75	0.0797
1.5	0.094	2.5	0.118
3.5	0.137	5.5	0.155
8.5	0.185	11.5	0.239
18.3	0.301	24.4	0.529
32.4	0.617	48.8	0.744
48.8	0.744	80.5	0.867
147	0.923	225	0.931
398	0.937	577	0.931
702	0.942		

2280300
LPE/BHT/AQ/30C/L2280600
LPE/BHT/AQ/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.25	0.00167	0.25	0.0102
0.5	0.00281	0.5	0.0172
1.2	0.00374	1.2	0.0229
2.4	0.00555	2.4	0.0265
7.28	0.00788	5.75	0.0379
27.9	0.0116	24	0.0465
48.8	0.0175	31.4	0.127
76.3	0.0223	52	0.157
193	0.0268	121	0.225
364	0.0447	172	0.407
491	0.0556	16	0.491
912	0.0729	1216	0.575
1225	0.106	337	0.362
1565	0.13	508	0.758
2331	0.15	625	0.833
3028	0.216	1056	0.903
3677	0.239	1369	0.906
4373	0.251	1729	0.899
5113	0.265	2475	0.855
5404	0.318	3220	0.878

BPE/BHT/CO/30C/L

2310300
BPE/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	1.0E-3	0.017	0.0125
0.033	0.00129	0.033	0.00857
0.067	0.00202	0.067	0.0134
0.133	0.00377	0.133	0.0219
0.25	0.00552	0.25	0.0283
0.5	0.00694	0.483	0.0557
1.2	0.00968	0.983	0.088
2.4	0.0226	2.4	0.0.147
4.8	0.0337	6.7.5	0.0.247
9.7	0.0374	24.1	0.0.329
19.3	0.0748	230.4	0.0.376
36.4	0.0867	48	0.0.672
53.3	0.257	125	0.0.81
82.1	0.397	220	0.0.804
103.3	0.425	292	0.0.804
120.7	0.531	565	0.0.892
201.7	0.619	893	0.0.81
201.7	0.659	1924	0.0.776
	0.697	1924	0.0.775
	0.696		

BPE/BHT/EN/30C/L 2320700 2320600 BPE/BHT/EN/60C/L1 2320602 BPE/BHT/EN/60C/L2

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0. 017	0. 00187	0. 017	0. 00779	0. 017	0. 00603	0. 00603	0. 00603
0. 033	0. 00255	0. 033	0. 00937	0. 033	0. 00763	0. 00763	0. 00763
0. 067	0. 00355	0. 067	0. 0166	0. 067	0. 0111	0. 0111	0. 0111
0. 133	0. 00526	0. 133	0. 0232	0. 133	0. 0169	0. 0169	0. 0169
0. 25	0. 00685	0. 25	0. 0343	0. 25	0. 0265	0. 0265	0. 0265
0. 5	0. 00943	0. 5	0. 0584	0. 5	0. 0437	0. 0437	0. 0437
1. 5	0. 0137	1. 5	0. 137	1. 5	0. 0766	0. 0766	0. 0766
2. 5	0. 0226	2. 5	0. 186	2. 5	0. 106	0. 106	0. 106
4	0. 0303	4	0. 233	4	0. 2295	0. 2295	0. 2295
6	0. 0398	6	0. 291	6	0. 2356	0. 2356	0. 2356
7	0. 0438	7	0. 389	7	0. 25	0. 25	0. 25
24	0. 0397	24	0. 432	24	0. 563	0. 563	0. 563
48	0. 0161	48	0. 178	48	0. 614	0. 614	0. 614
54.	0. 0161	54.	0. 178	54.	0. 636	0. 636	0. 636
121	0. 0302	121	0. 302	121	0. 642	0. 642	0. 642
229	0. 0434	229	0. 434	229	0. 642	0. 642	0. 642
389	0. 0581	389	0. 581	389	0. 642	0. 642	0. 642
555	0. 0667	555	0. 667	555	0. 642	0. 642	0. 642
817	0. 0733	817	0. 733	817	0. 642	0. 642	0. 642
1037	0. 0774	1037	0. 774	1037	0. 642	0. 642	0. 642
1206	0. 0785	1206	0. 785	1206	0. 642	0. 642	0. 642
1565	0. 0796	1565	0. 796	1565	0. 642	0. 642	0. 642
2065	0. 0814	2065	0. 814	2065	0. 642	0. 642	0. 642
2716	0. 0817	2716	0. 817	2716	0. 642	0. 642	0. 642
2716	0. 0822	2716	0. 822	2716	0. 642	0. 642	0. 642

	BPE/BHT/HP/300C/U	BPE/BHT/HP/600C/U1	BPE/BHT/HP/600C/U2	BPE/BHT/HP/600C/U3
t, h	Mt/Mo	t, h	Mt/Mo	t, h
0.017	0.00947	0.017	0.0548	0.017
0.033	0.015	0.033	0.0918	0.033
0.067	0.0244	0.067	0.153	0.067
0.117	0.0364	0.117	0.231	0.115
0.183	0.0492	0.183	0.318	0.25
0.25	0.0627	0.25	0.401	0.5
0.5	0.0108	0.5	0.661	1
0.75	0.151	0.783	0.779	2.2
1	0.0192	1	0.803	4
1.5	0.0235	1.5	0.822	7
2.5	0.0359	2.5	0.824	7
4	0.0449	4	0.825	7
5.5	0.065	5.5	0.826	7
8	0.741	8	0.826	7
24.1	0.792	23.8	0.828	7
48	0.812	31.9	0.828	7
80.9	0.814	47.9	0.828	7
249	0.815	80.	0.829	7
384	0.816	144	0.831	7
557	0.817	225	0.832	7
		388	0.833	7
		395	0.834	7
		702		

2370300
BPE/BHT/T0/30C/L2370600
BPE/BHT/T0/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 017	0. 00103	0. 017	0. 00764
0. 033	0. 00115	0. 033	0. 00889
0. 067	0. 00228	0. 067	0. 0145
0. 133	0. 00284	0. 133	0. 0217
0. 25	0. 00391	0. 25	0. 0365
0. 5	0. 00565	0. 5	0. 061
1. 03	0. 00809	1. 03	0. 0995
1. 5	0. 0104	1. 5	0. 16
2. 5	0. 0142	2. 5	0. 265
4	0. 0249	4	0. 349
6	0. 0294	6	0. 403
7	0. 0349	7	0. 67
11	0. 0711	11	0. 701
12	0. 124	12	0. 725
14	0. 255	14	0. 732
24	0. 443	24	0. 747
31	0. 582	31	0. 751
48	0. 658	48	0. 749
125	0. 704	125	0. 751
313	0. 731	313	0. 746
533	0. 731	533	0. 748
793	0. 731	793	0. 773
1237	0. 731	1237	0. 76
1561	0. 731	1561	0. 762
2524	0. 731	2524	0. 762
2524	0. 731	2524	0. 762

2380600
BPE/BHT/AQ/600C/L

t, h	Mt/Mo
1	0.0137
2.07	0.0195
4	0.0263
6.5	0.0346
24.4	0.0602
97.8	0.088
373	0.141
701	0.211
1970	0.378
2000	0.412
2000	0.417
2717	0.508

2430601
BPE/EHT/HP/60C/U1

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0238	0.017	0.0253
0.033	0.0412	0.033	0.0441
0.067	0.0755	0.067	0.0816
0.133	0.101	0.133	0.111
0.25	0.109	0.25	0.118
0.5	0.112	0.5	0.122
1.02	0.115	1	0.124
2.4	0.116	2	0.126
6.08	0.118	4.25	0.127
8	0.119	6	0.128
24.5	0.121	7.27	0.129
25.7	0.121	24.1	0.13

2430602
BPE/EHT/HP/60C/U2

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	0.017	0.0253
0.033	0.033	0.033	0.0441
0.067	0.067	0.067	0.0816
0.133	0.133	0.133	0.111
0.25	0.25	0.25	0.118
0.5	0.5	0.5	0.122
1	1	1	0.124
2	2	2	0.126
4	4	4.25	0.127
6	6	6	0.128
7.27	7.27	7.27	0.129
24.1	24.1	24.1	0.13
25	25	25	0.13

2510300
BPE/BHT/CO/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00611	1.5	0.0862
0.033	0.00915	2.5	0.113
0.067	0.0163	3.5	0.139
0.133	0.0236	5.5	0.179
0.25	0.0342	8.5	0.227
0.5	0.0475	24.9	0.419
1.5	0.0598	32.3	0.479
2.5	0.0698	48.4	0.593
3.5	0.0862	80.3	0.74
5.5	0.113	144	0.887
8.5	0.139	221	0.964
24.9	0.179	393	0.998
32.3	0.227	577	1.01
48.4	0.419	866	1.01
80.3	0.479	1207	0.989

2510600
BPE/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.017	0.017	0.0361
0.033	0.033	0.067	0.0501
0.067	0.067	0.117	0.0722
0.133	0.117	0.183	0.0956
0.25	0.25	0.75	0.119
0.5	0.5	0.75	0.152
1.5	0.5	0.75	0.253
2.5	0.5	0.75	0.334
5.5	0.5	1.5	0.401
8.5	0.5	5.5	0.519
24.9	0.5	24.9	0.613
32.3	0.5	32.3	0.691
48.4	0.5	8.5	0.805
80.3	0.5	5.5	0.926
144	0.5	8.5	0.987
221	0.5	24.9	0.998
393	0.5	32.3	1.01
577	0.5	48.4	1.01
866	0.5	56.4	1.01
1207	0.5	96.4	1.01
		294	0.993

2520300
BPE/BHT/EH/30C/L2520600
BPE/BHT/EN/60C/L12520602
BPE/BHT/EN/60C/L2

t, h	Mt/Mo						
0.017	0.0105	0.017	0.0534	0.017	0.042	0.017	0.042
0.033	0.0157	0.033	0.0675	0.033	0.0578	0.067	0.0578
0.067	0.0235	0.067	0.0914	0.067	0.0749	0.117	0.0997
0.117	0.0313	0.117	0.118	0.117	0.132	0.183	0.132
0.183	0.0378	0.183	0.153	0.183	0.183	0.183	0.183
0.25	0.0441	0.25	0.05	0.25	0.159	0.25	0.159
0.5	0.0618	0.5	0.05	0.5	0.277	0.5	0.277
0.75	0.074	0.75	0.05	0.75	0.363	0.75	0.363
1.5	0.0855	1.5	0.06	1.5	0.44	1.5	0.44
2.5	0.106	2.5	0.122	2.5	0.57	2.5	0.57
4	0.129	4	0.139	4	0.66	4	0.66
5.5	0.177	5.5	0.211	5.5	0.844	5.5	0.844
8	0.259	8	0.259	8	0.945	8	0.945
25.8	0.488	24	0.488	24	0.47	24	0.47
38.2	0.599	32	0.599	32	0.47	32	0.47
55.2	0.709	55	0.709	55	0.56	55	0.56
148	0.931	120	0.931	120	0.56	120	0.56
288	1	216	1	216	1	123	1
384	0.992	361	0.992	361	0.994	361	0.994
649	1	650	1	650	0.994	650	0.994
866	0.996	866	0.996	866	0.996	866	0.996
1177	1	1177	1	1177	1	1177	1
1566	0.998	1566	0.998	1566	0.998	1566	0.998
2022	0.99	2022	0.99	2022	0.99	2022	0.99

2595300
BPE/BHT/.95EN/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.00222	0.067	0.00751
0.133	0.0118	0.25	0.021
0.5	0.0286	1	0.0435
2	0.0658	3.97	0.103
5.48	0.13	24.4	0.324
30.1	0.384	53.2	0.523
125	0.744	292	0.877
480	0.919	649	0.915
700	0.93		

2595600
BPE/BHT/.95EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.033	0.017	0.067	0.033
0.133	0.133	0.25	0.25
1	0.5	3	1.95
4	4	24.	6.75
30.	24.	30.	24.
96.	96.	145	145

2530301
BPE/BHT/HP/30C/U

2530601
BPE/BHT/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0538	0.017	0.24
0.033	0.0797	0.033	0.382
0.067	0.127	0.067	0.616
0.117	0.192	0.133	0.9
0.183	0.273	0.25	0.961
0.25	0.353	0.5	0.964
0.5	0.683	1	0.965
1	0.942	2	0.965
1.5	0.982	4	0.965
2.5	0.989	5	0.965
4	0.991	7.5	0.965
5	0.992	24.3	0.965
5.5	0.993	52.2	0.965
8.1	0.993		
24	0.993		
32	0.993		
53.9	0.993		

BPE/BHT/ON/30C/L

BPE/BHT/ON/60C/L1

BPE/BHT/ON/60C/L2

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00479	0.017	0.017	0.033	0.0582	0.017	0.0275
0.033	0.00718	0.033	0.0754	0.067	0.033	0.041	0.041
0.067	0.0114	0.067	0.106	0.117	0.067	0.0566	0.0566
0.117	0.0179	0.133	0.151	0.183	0.111	0.077	0.077
0.183	0.021	0.25	0.218	0.316	0.175	0.111	0.111
0.2	0.029	0.5	0.551	1.4	0.557	0.279	0.279
0.5	0.0394	1	0.944	6	0.64	0.434	0.434
0.833	0.0579	1.88	0.771	23.9	0.986	0.986	0.986
1.5	0.0833	4	0.944	229.9	0.996	0.996	0.996
2.5	0.123	19	0.64	51.5	1.12	1.0	1.0
4.27	0.25	51.3	0.982	72.3	0.968	0.998	0.998
8.67	0.19	603	0.999	145	0.965	0.997	0.997
23.8	0.513	1	0.999	216	0.962	0.998	0.998
31.1	0.603	1.1	0.999	385	0.969	0.998	0.998
55.1	0.799	120	0.936	1182	0.982	0.982	0.982
120	0.936	216	0.966	865	0.972	0.972	0.972
216	0.966	385	0.974	554	0.982	0.982	0.982
385	0.974	1182	0.982	1182	0.982	0.982	0.982

2560300
BPE/BHT/TB/30C/L2560600
BPE/BHT/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.067	0.0225	0.017	0.042
0.133	0.0332	0.033	0.0587
0.25	0.0432	0.067	0.0841
0.5	0.0599	0.133	0.12
1	0.0842	0.25	0.175
2	0.118	0.5	0.274
4	0.17	1	0.424
6	0.214	2	0.634
7, 42	0.24	4	0.837
24	0.35	6	0.99
38, 6	0.499	7, 33	1.01
48, 2	0.609	24	0.952
121	0.84	48	0.950
216	0.939	97	0.937
265	0.965	192	0.935
360	0.971	313	0.934
481	0.971	605	0.933
793	0.977		

2580300
BPE/BHT/AQ/30C/L

t, h	Mr/Mo
0.25	0.0085
0.5	0.0106
1	0.0158
2	0.0202
4	0.0272
23.5	0.0433
96.6	0.0728
192	0.107
364	0.161
697	0.272
1442	0.4
2133	0.467
2655	0.473
3460	0.425
3768	0.548

2580600
BPE/BHT/AQ/60C/L

t, h	Mr/Mo
0.25	0.083
0.5	0.167
1	0.25
2	0.0216
4	0.0317
7.42	0.0419
24.6	0.0876
72.3	0.155
244	0.199
601	0.464
1347	0.62
2693	0.795
3365	0.849
3580	0.867
3	0.904

2590400
BPE/BHT/HB/40C/L12590402
BPE/BHT/HB/40C/L22590600
BPE/BHT/HB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00702	0.017	0.00648	0.017	0.00648	0.017	0.03
0.033	0.0188	0.033	0.0145	0.033	0.0145	0.033	0.0579
0.067	0.0206	0.067	0.021	0.067	0.021	0.067	0.0774
0.133	0.0274	0.133	0.0333	0.133	0.0333	0.133	0.129
0.267	0.0534	0.25	0.0488	0.25	0.0488	0.25	0.197
0.5	0.0747	0.5	0.0796	0.5	0.0796	0.5	0.296
1	0.118	1	0.121	1	0.121	1	0.462
2.42	0.213	1.92	0.182	1.97	0.182	1.97	0.659
4	0.289	1.4	0.293	4	0.293	4	0.857
5.45	0.356	6.05	0.381	5.98	0.381	5.98	0.93
6.75	0.437	7.42	0.428	7.42	0.428	7.42	0.974
8.95	0.52	24	0.787	24	0.787	24	0.969
11.3	0.52	30	0.87	30	0.87	30	0.976
32	0.874	49	0.925	49	0.925	49	0.984
57.9	0.966	124	0.979	124	0.979	124	0.973
126	0.99	318	1	318	1	318	0.958
222	0.985	529	0.979	529	0.979	529	0.958
297	0.993	630	0.984	630	0.984	630	0.958

2610600
LPE/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00208	124	6.75
0.033	0.00358	231.4	
0.067	0.00469	76	
0.133	0.0132	52	
0.25	0.0271	145	
0.5	0.0428	240	
1	0.0779	361	
2	0.131	532	
4	0.202	556	
6	0.256	556	
8	0.298	556	
14	0.548	145	
24	0.658	240	
31.5	0.809	361	
46	0.913	532	
75	0.958	556	
124	0.966	556	
148	0.945	145	
172	0.929	240	
265	0.925	361	
317	0.93	481	
417	0.905	652	
673	0.905	673	
100	0.905		
124	0.917		
148	0.93		
172	0.929		
265	0.925		
317	0.93		
417	0.905		
673	0.905		

2620600
LPE/BHT/EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00502	0.017	0.00502
0.033	0.00704	0.033	0.00704
0.067	0.0118	0.067	0.0118
0.133	0.0187	0.133	0.0187
0.25	0.0296	0.25	0.0296
0.5	0.0519	0.5	0.0519
1	0.0857	1	0.0857
2	0.139	2	0.139
4	0.224	4	0.224
6	0.299	6	0.299
14	0.378	14	0.378
24	0.538	24	0.538
31.5	0.763	31.5	0.763
46	0.905	46	0.905
75	0.917	75	0.917
124	0.93	124	0.93
148	0.93	148	0.93
172	0.929	172	0.929
265	0.925	265	0.925
317	0.93	317	0.93
417	0.905	417	0.905
673	0.905	673	0.905

LPE/BHT/HP/30C/U

2630301
LPE/BHT/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0226	0.017	0.0998
0.033	0.0392	0.033	0.166
0.067	0.0682	0.067	0.274
0.117	0.104	0.133	0.464
0.183	0.148	0.25	0.759
0.25	0.183	0.5	0.94
0.75	0.334	1.24	0.967
1.5	0.495	2.4	0.971
2.5	0.635	5	0.971
5	0.888	6	0.971
5	0.989	7	0.971
8	0.996	24	0.972
11	0.999	25	0.972
15	0.999		

2710300
PP/BHT/CO/30C/L

2710600
PP/BHT/CO/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
7.5	0.00303	0.5	0.00566
25	0.00838	0.033	0.00566
31.5	0.00845	0.067	0.00918
57.1	0.0155	0.133	0.0304
97.4	0.0219	0.25	0.0223
194	0.0354	0.5	0.0409
366	0.059	1	0.0636
605	0.079	2	0.0951
871	0.101	4	0.144
1181	0.122	6	0.13
1538	0.14	7	0.52
1946	0.164	8	0.202
2379	0.184	9	0.368
2907	0.208	10	0.429
3484	0.231	48	0.535
4081	0.243	55	0.574
4731	0.28	72.1	0.652
5426	0.301	144	0.814
		244	0.844
		385	0.833
		529	0.867
		844	0.864

2720300
PP/BHT/EN/30C/L2720600
PP/BHT/EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
1.5	0.00317	0.5	0.00401
2.5	0.00356	0.5	0.00481
3.5	0.00808	0.5	0.0108
5.5	0.0147	0.5	0.0177
6.77	0.0398	0.5	0.0393
8.83	0.0641	0.5	0.0641
25.4	0.0919	0.5	0.0919
34.6	0.116	0.5	0.116
50.5	0.168	0.5	0.168
84.4	0.222	0.5	0.222
121	0.278	0.5	0.278
223	0.326	0.5	0.326
386	0.38	0.5	0.38
625	0.437	0.5	0.437
869	0.484	0.5	0.484
1181	0.526	0.5	0.526
1564	0.563	0.5	0.563
1899	0.589	0.5	0.589
2407	0.634	0.5	0.634
2911	0.671	0.5	0.671
3411	0.692	0.5	0.692
4156	0.706	0.5	0.706
4880	0.730	0.5	0.730
5595	0.754	0.5	0.754
6340	0.787	0.5	0.787

2736301
PP/BHT/HP/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.021	0.015	0.015	0.017
0.033	0.0213	0.033	0.033
0.067	0.0352	0.067	0.067
0.117	0.0508	0.133	0.133
0.183	0.0668	0.25	0.25
0.35	0.0803	0.5	0.5
0.65	0.0983	1.5	1.5
1.42	0.123	2.35	2.35
2.75	0.161	5.5	5.5
3.5	0.202	7.5	7.5
4.67	0.295	24.1	24.1
6.67	0.775	51.9	51.9
8.88	0.865	88.7	88.7
24.1	0.872	0.884	0.884
54.1	0.879	0.881	0.881
		0.885	0.885
		0.886	0.886
		0.887	0.887
		0.888	0.888

2730601
PP/BHT/HP/60C/U

t, h	Mt/Mo
0.017	0.017
0.033	0.033
0.067	0.067
0.133	0.133
0.25	0.25
0.5	0.5
1.5	1.5
2.35	2.35
5.5	5.5
7.5	7.5
24.1	24.1
51.9	51.9
88.7	88.7
0.884	0.884
0.881	0.881
0.884	0.884
0.881	0.881
0.885	0.885
0.885	0.885
0.886	0.886
0.887	0.887
0.887	0.887
0.888	0.888

2760300
PP/BHT/TB/30C/L2760600
PP/BHT/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
1.5	7.89E-4	0.067	0.00587
2.5	0.0114	0.133	0.00787
3.5	0.00394	0.25	0.0152
5.5	0.00306	0.5	0.0235
7.5	0.00479	1.17	0.0443
12.5	0.0051	1.95	0.071
24.2	0.0109	4.25	0.113
41.2	0.0326	6.2	0.187
73	0.0539	8.8	0.232
125	0.0794	11.5	0.365
242	0.126	14.2	0.555
412	0.151	17.5	0.648
606	0.184	21.7	0.837
875	0.223	24.7	0.842
1184	0.267	27.7	0.856
1516	0.309	31.7	0.867
1922	0.361	37.2	0.882
2402	0.407	42.2	
2839	0.457	47.2	
3339	0.505	52.2	
4084	0.587	57.2	
4808	0.65	62.2	
5547	0.702	67.2	
6222	0.752	72.2	
6992	0.717		

2780300
PP/BHT/AQ/30C/L

t, h Mt/Mo

23.2
51.5
94.3
176
1293
432
602
8625
1154
1154
2285
3604
4274
4997
5736
6411
7231

0.00434
0.00819
0.0145
0.0206
0.0354
0.0449
0.0545
0.0647
0.076
0.0947
0.11
0.124
0.136
0.145
0.145
0.158
0.152
0.209

2780600
PP/BHT/AQ/60C/L

t, h Mt/Mo

0.25
0.5
1
2
4.12
6.33
24
293.2
96.5
176
337
337
505
673
865
1204
1537
1873
2113
2449
2617
2788

0.00254
0.00312
0.0061
0.0117
0.0177
0.0243
0.05
0.0878
0.145
0.205
0.314
0.384
0.447
0.506
0.554
0.586
0.596
0.595
0.588
0.583

2810300
PP/C18/C0/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0024	0.017	0.0172
0.033	0.00301	0.033	0.0246
0.067	0.00457	0.067	0.0425
0.117	0.00585	0.133	0.048
0.183	0.00761	0.25	0.0945
0.25	0.00904	0.5	1.0
0.417	0.0116	1	2
0.667	0.0156	2	4
1.1	0.025	4	6
1.5	0.025	6	7.5
2.2	0.025	7.5	24.3
2.7	0.025	7.5	31.5
4.5	0.0535	14.5	48.8
5.5	0.0535	36.1	1.0
6.7	0.0653		0.999
7.5	0.0653		
7.5	0.131		
7.5	0.151		
7.5	0.229		
14.6	0.309		
24.0	0.388		
40.9	0.491		
60.1	0.569		
84.7	0.646		
117.9	0.721		
151.4	0.779		
192.5	0.834		
240.2	0.878		
307.7	0.924		
373.7	0.951		
447.1	0.976		
516.3	0.986		
588.5	0.996		
662.5	0.992		

PP/C18/EN/300C/L

2820300
PP/C18/EN/600C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00383	0.017	0.017
0.033	0.00551	0.033	0.03
0.067	0.00774	0.067	0.0476
0.117	0.0109	0.133	0.0781
0.183	0.0134	0.25	0.123
0.25	0.0166	0.5	0.195
0.417	0.0242	1	0.308
0.667	0.0332	2	0.48
1.5	0.0425	4	0.7
2.5	0.0542	6.25	0.833
5.5	0.0641	23.4	0.949
7.5	0.0737	271.9	0.957
7.7	0.0966	121	0.955
7.68	0.118	193	0.936
25.9	0.272	289	0.941
32	0.304	385	0.996
79.2	0.486		
127	0.61		
243	0.794		
408	0.908		
602	0.964		
873	0.988		
1184	1		
1517	1		
1926	1		
1999	0.993		

2830391
PP/C18/HP/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0378	0.017	0.128
0.033	0.0573	0.033	0.184
0.067	0.0853	0.067	0.304
0.117	0.116	0.133	0.708
0.183	0.15	0.25	0.997
0.25	0.18	0.5	1
0.35	0.223	1	1
0.5	0.287	2	1
0.75	0.426	22.8	1
1.0	0.908	51.8	1
1.25	0.995		
1.5	0.999		
1.75	0.999		
2.25	1		
2.5	1		
2.75	1		
3.5	1		
3.98	1		
5	1		
6.67	1		
8.83	1		
24	1		
31.8	1		

2830601
PP/C18/HP/60C/U

t, h	Mt/Mo
0.017	0.0378
0.033	0.0573
0.067	0.0853
0.117	0.116
0.183	0.15
0.25	0.18
0.35	0.223
0.5	0.287
0.75	0.426
1.0	0.908
1.25	0.995
1.5	0.999
1.75	0.999

2860300
PP/C18/TB/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00207	0.017	0.0206
0.033	0.00385	0.033	0.029
0.067	0.00607	0.067	0.0445
0.117	0.00808	0.133	0.064
0.183	0.0104	0.225	0.0958
0.25	0.0127	0.5	0.155
0.417	0.0172	1.5	0.223
0.667	0.0224	2.4	0.332
1.1	0.0288	3.3	0.486
1.5	0.0356	5.5	0.597
2.5	0.0409	6.7	0.668
3.3	0.0464	7.27	0.699
5.27	0.0578	7.33	0.695
7.33	0.0699	15.8	0.805
11.5	0.0785	18.1	0.885
17.1	0.085	22.8	0.905
24.1	0.098	24.6	0.926
31.2	0.101	38.1	0.925
47.4	0.128	46.1	0.905
55.2	0.144	59.9	0.905
144	0.144	60.8	0.905
218	0.144	66.8	0.905
38.8	0.1516	118.0	0.954
60.8	0.1516	190.2	0.954
86.8	0.1516	240.6	0.99
118.0	0.1516	290.5	0.99
190.2	0.1516	340.9	0.993
240.6	0.1516	468.1	0.996

2880300
PP/C18/AQ/30C/L

2880600
PP/C18/AQ/60C/L

t, h Mt/Mo

T₀
9173 0.0005±0.0002

t, h	Mt/Mo	t, h	Mt/Mo
27.8	6.7E-4	96.6	7.53E-4
169.	6.26E-4	1761	6.89E-4
77.00	6.89E-4	1769	6.0103
12092	6.011	2113	6.0115
22833	8.92E-4	3941	7.65E-4
4632	8.76E-4	5405	9.7E-4
5405	9.7E-4	6072	8.41E-4
6865	8.65E-4	7657	8.42E-4
8760	9.41E-4	9582	9.72E-4

2910300
 PP/C32/C0/30C/L
 t, h Mt/Mo

0.017	0.0394
0.033	0.0419
0.067	0.0487
0.117	0.0499
0.183	0.0496
0.25	0.0496
0.33	0.0496
0.5	0.0496
1.247	0.0508
2.47	0.0519
5.26	0.0525
9.67	0.0535
29.1	0.0555
52.6	0.0565
96.7	0.0576
291	0.0668
337	0.0768
577	0.0876
840	0.0976
1157	0.1029
1513	0.129
1922	0.156
2355	0.196
2790	0.231
3462	0.264
4057	0.339
4710	0.408
5401	0.453
6144	0.491
6918	0.538
7274	0.579
	0.627
	0.691

2910600
 PP/C32/C0/60C/L
 t, h Nb/Mo

0.017	0.0374
0.033	0.0406
0.067	0.0434
0.117	0.0456
0.183	0.0502
0.25	0.0556
0.417	0.0714
1.15	0.0714
1.42	0.14
1.92	0.166
2.53	0.193
3.53	0.234
5.53	0.282
7.27	0.342
11.66	0.637
25.7	0.867
52.1	0.867
82.8	0.968
125	0.994
245	0.994
366	0.995

2920300
PP/C32/EH/30C/L

2920600
PP/C32/EH/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0136	0.017	0.0136
0.033	0.0243	0.033	0.0243
0.067	0.0341	0.067	0.0341
0.133	0.0356	0.117	0.0421
0.25	0.0362	0.183	0.0483
0.5	0.0382	0.25	0.059
1.24	0.0411	0.417	0.0929
2.4	0.0497	0.667	0.132
6.7	0.052	1.1	0.172
12.4	0.0594	1.2	0.219
24.8	0.0921	2.3	0.274
49.7	0.118	3.5	0.349
99.4	0.138	5.2	0.394
198.1	0.201	7.25	0.539
396.2	0.239	9.25	0.646
792.4	0.279	11.25	0.942
1584.8	0.313	13.25	0.955
3169.6	0.36	15.25	0.972
6339.2	0.412	17.25	0.972
1269.1	0.414	19.25	0.981
2538.2	0.481	21.25	0.981
5076.4	0.533	23.25	0.981
10152.8	0.587	25.25	0.981
20305.6	0.625	27.25	0.981
40611.2	0.653	29.25	0.981
81222.4	0.687	31.25	0.981
16244.8	0.722	33.25	0.981
32489.6	0.74	35.25	0.981
64979.2	0.751	37.25	0.981
12995.8	0.768	39.25	0.981
25991.6	0.768	41.25	0.981
51983.2	0.768	43.25	0.981
103966.4	0.768	45.25	0.981
207932.8	0.768	47.25	0.981
415865.6	0.768	49.25	0.981
831731.2	0.768	51.25	0.981
166344.8	0.768	53.25	0.981
332689.6	0.768	55.25	0.981
665379.2	0.768	57.25	0.981
133079.6	0.768	59.25	0.981
266159.2	0.768	61.25	0.981

2930301
PP/C32/HP/30C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0354	1.1.97	
0.033	0.0415	4.5.67	
0.067	0.0507	24.1	
0.15	0.0708	31.6	
0.25	0.0902	121	1.96.5
0.5	0.131		
	0.216		
	0.813		
	0.996		
	0.989		
	0.991		
	0.993		
	0.995		
	0.995		

2930601
PP/C32/HP/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0.008	0.0673	0.017	0.0906
0.033	0.132	0.067	0.207
0.1	0.282	0.1	0.401
0.2	0.644	0.167	0.644
0.267	0.814	0.133	0.906
0.333	0.953	0.233	0.975
0.333	0.975	0.533	0.986
0.333	0.986	0.833	0.995
0.333	0.997	1.33	0.997
2.3	0.997	2.83	0.998
3.9	0.998	19.1	0.998
127.9	0.998	51.9	0.998
73.7	0.998	93.7	0.998

2960300
PP/C32/TB/30C/L

2960600
PP/C32/TB/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0255	0.017	0.0404
0.033	0.0389	0.033	0.0415
0.067	0.0521	0.067	0.0436
0.117	0.0534	0.117	0.0475
0.183	0.0551	0.0.233	0.0569
0.3	0.0543	0.0.533	0.0904
0.5	0.0545	0.0.833	0.1165
1.2	0.0553	1.0.33	0.155
2.4	0.0575	2.0.833	0.202
7.7	0.0592	3.0.25	0.361
24	0.0626	4.0.75	0.419
31	0.0651	7.0.25	0.92
121	0.0817	100	0.983
217	0.101	100	0.977
390	0.138	221	0.988
628	0.182	413	0.991
865	0.246		
1217	0.307		
173	0.358		
1540	0.426		
1540	0.504		
1671	0.614		
2402	0.674		
2931	0.753		
3507	0.825		
4085	0.879		
4730	0.929		
5426	0.945		
6196	0.957		
6964	0.958		
7374	0.957		

2980600
PP/C32/AQ/60C/L

t, h Ht/Mo

T0 0.0005±0.0002
8065

2980300
PP/C32/AQ/30C/L

t, h Mt/Mo

t, h	Mt/Mo
4705	5.05E-4
5401	9.73E-4
6125	0.00291
6937	0.00507
8068	0.0085
	0.0164

3040301
PP/C18/0D/30C/U3040601
PP/C18/0D/60C/U

t, h	Mt/Mo	t, h	Mt/Mo
0. 005	0. 0294	0. 005	0. 0005
0. 011	0. 0639	0. 011	0. 639
0. 017	0. 0785	0. 017	0. 661
0. 025	0. 0975	0. 025	0. 672
0. 033	0. 114	0. 033	0. 694
0. 05	0. 136	0. 05	0. 696
0. 067	0. 157	0. 067	0. 714
0. 1	0. 188	0. 1	0. 73
0. 133	0. 215	0. 133	0. 759
0. 167	0. 238	0. 167	0. 785
0. 25	0. 239	0. 25	0. 808
0. 333	0. 232	0. 333	0. 856
0. 5	0. 406	0. 5	0. 891
0. 75	0. 499	0. 75	0. 938
1. 02	0. 584	1. 02	0. 972
1. 5	0. 701	1. 5	0. 988
2	0. 786	2	0. 998
3	0. 89	3	1
4. 5	0. 959	4. 5	1
6. 25	0. 987	6. 25	1
8. 93	0. 997	8. 93	1
26. 1		26. 1	
30. 9		30. 9	

E. 05uA/BHT/CO/30C/L
3110300

E. 05uA/BHT/CO/60C/L
3110600

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.00458	0.017	0.0668
0.033	0.0478	0.033	0.103
0.067	0.0208	0.067	0.155
0.133	0.0253	0.133	0.234
0.25	0.0342	0.25	0.292
0.5	0.0913	0.5	0.478
1.1	0.128	1.1	0.688
1.92	0.207	1.98	0.824
3.95	0.305	3.93	0.865
7.85	0.369	5.5	0.878
23.6	0.426	6.83	0.873
23.7	0.713	24.1	0.867
48.3	0.778	121	0.877
54.8	0.818		
77.5	0.839		
144	0.868		
222	0.883		
341	0.881		
384	0.882		
	0.88		

E. 05U/A/BHT/EN/30C/L
3120300

E. 05U/A/BHT/EN/60C/L
3120600

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0209	0.017	0.0824
0.033	0.03	0.033	0.129
0.067	0.0413	0.067	0.182
0.133	0.0629	0.133	0.262
0.25	0.0971	0.25	0.382
0.5	0.127	0.5	0.568
1.02	0.186	1	1.98
1.97	0.263	1.97	3.98
4.42	0.392	4.42	5.52
24.1	0.471	24.1	6.78
30.8	0.845	30.8	8.64
36.3	0.877	36.3	8.88
151	0.888	151	8.92
264	0.892	264	8.92
461	0.89	461	8.78
484	0.893	484	8.74
152	0.893	152	8.7
172	0.893	172	8.93

E. 05uA/BHT/.95EN/60C/L

3195600
E. 05uA/BHT/.95EN/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.9906	0.017	0.0291
0.033	0.127	0.033	0.0391
0.067	0.173	0.067	0.0521
0.117	0.223	0.117	0.0688
0.183	0.284	0.183	0.0845
0.283	0.369	0.283	0.104
0.433	0.48	0.433	0.127
0.667	0.618	0.667	0.155
1.1	0.741	1.1	0.19
1.5	0.84	1.5	0.235
2.2	0.88	2.2	0.269
4.2	0.905	4.25	0.394
6.52	0.921	6	0.473
9.2	0.921	8.82	0.573
22.5	0.919	22.5	0.798
32.4	0.922	32.5	0.853
51.8	0.913	48.2	0.889

t, h	Mt/Mo
83.3	0.911
145	0.916
217	0.925
384	0.918
581	0.909

3130300
E. 05VA/BHT/HP/30C/L

3130301
E. 05VA/BHT/HP/30C/U

3130600
E. 05VA/BHT/HP/60C/L

t, h	Mt/Mo
0.017	0.115
0.033	0.171
0.067	0.272
0.133	0.475
0.25	0.758
0.5	0.883
1.25	0.904
3	0.907
4.3	0.902
6.3	0.913
23.2	0.893
30.2	0.893
48.1	0.892
120	0.891
169	0.894

t, h	Mt/Mo
0.006	0.0472
0.011	0.0721
0.017	0.0938
0.025	0.121
0.033	0.145
0.067	0.187
0.067	0.224
0.15	0.292
0.2	0.386
0.45	0.479
1.5	0.646
3	0.786
4.5	0.853
6.67	0.875
14	0.911
23	0.921
30	0.921
48	0.925
120	0.933
169	0.941

3180300
E. 05VA/BHT/AQ/30C/L

3180600
E. 05VA/BHT/AQ/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 033	0. 00503	0. 033	0. 0152
0. 083	0. 0074	0. 083	0. 0212
0. 167	0. 00891	0. 167	0. 0274
0. 333	0. 0122	0. 25	0. 0337
0. 583	0. 0142	1	0. 05
1. 47	0. 0219	3	0. 0792
3. 67	0. 036	5. 07	0. 092
6. 62	0. 0515	7. 12	0. 102
24. 4	0. 0915	24. 2	0. 122
54	0. 122	30. 9	0. 132
149	0. 151	53. 5	0. 151
3313	0. 163	120	0. 204
5556	0. 183	198	0. 257
817	0. 2	293	0. 319
1033	0. 22	457	0. 384
1321	0. 226	700	0. 49
1657	0. 236	961	0. 565
2164	0. 257	1177	0. 61
2763	0. 286	1465	0. 656
3409	0. 307	1801	0. 67
3845	0. 343	2308	0. 665
		2907	0. 688
		3479	0. 696
		3966	0. 708

3210300
E. 13VA/BHT/CO/300C/L

3210690
E. 13VA/BHT/CO/600C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 017	0. 00949	0. 017	0. 017
0. 033	0. 0123	0. 033	0. 033
0. 067	0. 0209	0. 067	0. 067
0. 117	0. 0297	0. 117	0. 107
0. 183	0. 0364	0. 183	0. 136
0. 3	0. 0419	0. 283	0. 18
0. 5	0. 0652	0. 433	0. 237
0. 833	0. 0891	0. 667	0. 319
1. 33	0. 112	1. 5	0. 413
2. 3	0. 14	2. 3	0. 527
4. 5	0. 178	4. 25	0. 627
6. 25	0. 221	6. 25	0. 766
8. 5	0. 263	8. 5	0. 876
23. 8	0. 31	25. 3	0. 954
32. 4	0. 532	31. 9	1. 03
49. 6	0. 627	54. 7	1. 03
82	0. 756	78. 7	0. 99
121	0. 96	1	1
223	0. 989	651	0. 987
390	0. 997	870	
1183			

E. 13U/A/BHT/EH/30C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0178	0.017	0.0384
0.033	0.0257	0.033	0.0562
0.067	0.0362	0.067	0.0795
0.117	0.0492	0.117	0.12
0.183	0.062	0.183	0.16
0.25	0.0806	0.25	0.188
0.417	0.106	0.417	0.282
0.667	0.138	0.667	0.403
1.	0.191	1.	0.525
1.52	0.246	2.	0.682
2.	0.318	2.	0.835
2.5	0.468	2.	0.945
4.	0.59	3.	0.856
5.	0.75	5.	0.982
8.	0.91	5.	0.999
24.	0.975	5.	0.999
31.	0.977	7.	0.999
48.	0.992	7.	0.999
79.	0.998	31.	0.999
104	1.	1.	0.996
219	1.	1.	0.996
388	0.999	198	0.998
605	0.989	365	0.999
703	0.987	461	0.999
535	0.996	535	0.996

E. 13U/A/BHT/EH/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0178	0.017	0.0384
0.033	0.0257	0.033	0.0562
0.067	0.0362	0.067	0.0795
0.117	0.0492	0.117	0.12
0.183	0.062	0.183	0.16
0.25	0.0806	0.25	0.188
0.417	0.106	0.417	0.282
0.667	0.138	0.667	0.403
1.	0.191	1.	0.525
1.52	0.246	2.	0.682
2.	0.318	2.	0.835
2.5	0.468	5.	0.945
4.	0.59	5.	0.982
5.	0.75	7.	0.999
8.	0.91	7.	0.999
24.	0.975	31.	0.999
31.	0.977	48.	0.999
48.	0.992	79.	0.999
79.	0.998	104	1.
104	1.	219	1.
219	1.	388	0.999
388	0.999	605	0.989
605	0.989	703	0.987
703	0.987	461	0.999
461	0.999	535	0.996
535	0.996	535	0.996

E, 13VA/BHT/, .95EN/30C/L

3295300
3295600

t, h	Mt/Mo	t, h	Mt/Mo
0.017	0.0168	0.017	0.0528
0.033	0.0244	0.033	0.0745
0.067	0.0342	0.067	0.104
0.117	0.0462	0.117	0.137
0.183	0.0586	0.183	0.177
0.283	0.0734	0.283	0.231
0.433	0.0926	0.433	0.303
0.667	0.115	0.667	0.398
1.1	0.147	1.1	0.512
1.5	0.176	1.5	0.649
2.3	0.206	2.3	0.788
3.4	0.252	3.4	0.905
4.6	0.307	4.6	0.957
6.5	0.362	6.5	0.978
9.1	0.432	9.1	0.992
12.6	0.521	12.6	0.993
19.5	0.655	19.5	0.993
22.5	0.755	22.5	0.993
34.4	0.849	34.4	0.975
49.5	0.938	49.5	0.98
59.3	0.94	59.3	0.982
121	0.94	121	0.982
215	0.998	215	0.993
384	0.991	384	0.982
626	0.993	626	0.982
871	0.982	871	0.982

E. 13UA/BHT/HP/300C/U

3230301

t, h	Mt/Mo	t, h	Mt/Mo
0.005	0.0489	0.011	0.0729
0.017	0.0905	0.025	0.113
0.05	0.163	0.067	0.191
0.15	0.299	0.2	0.353
0.2	0.45	0.45	0.584
1.5	1.667	1.5	1.758
2.25	2.25	2.25	2.904
3.5	3.5	3.5	3.969
4.8	4.8	4.8	4.964
7	7	7	5.986
22	22	22	6.986
31	31	31	7.987
53.2	53.2	53.2	8.987
98.1	98.1	98.1	9.987
196	196	196	10.987

E. 13UA/BHT/HP/600C/U

3230601

t, h	Mt/Mo	t, h	Mt/Mo
0.005	0.005	0.011	0.126
0.017	0.017	0.025	0.191
0.025	0.025	0.033	0.243
0.05	0.05	0.067	0.308
0.1	0.1	0.167	0.365
0.2	0.2	0.233	0.463
0.45	0.45	0.55	0.55
0.584	0.584	0.703	0.703
0.904	0.904	0.916	0.916
1.758	1.758	2.233	0.932
2.904	2.904	3.5	0.956
4.964	4.964	4.333	0.975
6.986	6.986	6.83	0.983
7.987	7.987	8.5	0.986
8.987	8.987	9.05	0.987
9.987	9.987	10.3	0.988
10.987	10.987	11.6	0.988
12.3	12.3	12.3	0.989
14.05	14.05	14.05	0.989
16.69	16.69	16.69	0.989

3280300
E. 13UA/BHT/AQ/30C/L

3280600
E. 13UA/BHT/AQ/60C/L

t, h	Mt/Mo	t, h	Mt/Mo
0. 033	1. 83E-4	0. 083	5. 78E-4
0. 083	3. 95E-4	0. 167	0. 00148
0. 167	6. 09E-4	0. 333	0. 00172
0. 333	8. 37E-4	0. 667	0. 00284
0. 667	0. 00162	1. 5	0. 00373
1. 33	0. 00213	2. 25	0. 00456
2. 57	0. 00316	3. 75	0. 00764
4. 57	0. 00444	5. 25	0. 00974
7. 75	0. 00571	7. 5	0. 00952
21. 57	0. 00739	22. 1	0. 01372
23. 57	0. 0101	23. 5	0. 01522
25. 57	0. 0114	24. 5	0. 01599
27. 57	0. 0142	25. 2	0. 0342
29. 57	0. 0165	29. 2	0. 0491
31. 57	0. 0199	22. 3	0. 0664
33. 57	0. 0229	38. 5	0. 0998
35. 57	0. 0263	58. 2	0. 131
37. 57	0. 0305	87. 1	0. 176
39. 57	0. 0336	117. 6	0. 216
41. 57	0. 0398	153. 6	0. 265
43. 57	0. 0432	191. 9	0. 313
45. 57	0. 0496	240. 8	0. 371
47. 57	0. 0618	290. 4	
49. 57		341. 5	

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11. ABSTRACT (A 200-word or less factual summary of most significant information. If document includes a significant bibliography or literature survey, mention it here)				Food packaging is an inseparable part of modern life. Any substance that migrates from the packaging material into foods is viewed as indirect food additives. In connection with toxicological knowledge, it is important to know the amount of such indirect food additives expected to be present in the food during storage and processing. This program, sponsored by the Bureau of Foods of the Food and Drug Administration is to provide theoretical models, reliable data base, methodology to study the migration phenomena and to provide reasonable worst-case estimates for the concentrations of the indirect additives in food. In this final report we present the results of approximately 250 completed migration experiments based on radiotracer techniques on the migration of low molecular weight hydrocarbons and antioxidants from polyethylene, polypropylene and ethylene-vinyl acetate copolymers. Results of a study for the determination of relative diffusion coefficients of several probe molecules in the polyethylene melt by inverse gas chromatography are also presented. Based on these studies, ethanol appears to be a far more reasonable food-oil simulating solvent than n-heptane. Other pure or mixed triglycerides may also be considered as food-oil simulating solvents, however they may pose the same analytical difficulties as that of the food oil itself.		
12. KEY WORDS (Six to twelve entries; alphabetical order; capitalize only proper names; and separate key words by semicolons)						
antioxidants; diffusion; ethylene-vinyl acetate copolymers; food packaging; inverse gas chromatography; migration; oligomers; polyethylene; polypropylene; radiotracer						
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